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In vitro and in vivo chronopharmacology of a new generation of an organometallic anticancer drug complex

by

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A thesis submitted in the partial fulfilment of the requirements for the degree of

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‘Wissenschaft denkt nicht.’ Martin Heidegger
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**Declaration**

I declare that the work presented in this thesis with the title “*In vitro* and *in vivo* chronopharmacology of a new generation of an organometallic anticancer drug complex” is entirely researched and performed by myself unless stated differently in the text. The research in this thesis was done under the supervision of Professor Francis Lévi and Doctor Robert Dallmann. All sources of information have been acknowledged by references.

I further declare that none of this work has ever been submitted for any other degree.
The research presented in this thesis has highlighted the complementary and consistent links between in vitro and in vivo chronopharmacology findings, as shown for the anticancer drug candidate FY26. The use of 24-hour temperature schedules as an effective synchroniser of circadian clocks in vitro and that of core body temperature rhythm as a circadian biomarker in vivo has enabled the precise determination of the endogenous circadian times of best tolerability. This was achieved, as physiological rhythm had been introduced into cell culture conditions. Should these results be applicable to humans, least FY26 toxicity could occur at night, shortly after the physiological nadir in core body temperature. More generally, the synchronisation of healthy and cancer cells with 24-h periodic temperature schedules mimicking physiological cycles could have a high potential to identify optimal drug timing options and underlying chronopharmacokinetics and chronopharmacodynamics mechanisms, whilst reducing and replacing animal experiments, as illustrated here. This could indeed encourage the integration of chronotherapy concepts into drug development strategies. In such context, the objectives of animal chronotherapy studies might hence become mostly confirmatory. Thus, the use of core body temperature circadian cycle appears not only as crucial for the determination of the circadian timing resulting in best drug tolerability, antitumour efficacy, and quality of life, but it is also in agreement with the principles of the 3 R’s (reduction, refinement and replacement) and the public ethical opinion on animal experiments.
Extended Summary

Due to earth rotation, 24 h cycles with days and nights are generated on earth. In turn, the periodic availability of light resets endogenous circadian rhythm in plants, insects and mammals. Such rhythms are for example the daily changes of the sleep-wake cycle and core body temperature rhythms in mammals generated through a circadian timing system (CTS) which is synchronised by the day and night cycle.

The CTS consists of a central pacemaker in the suprachiasmatic nuclei coordinating rhythms in peripheral clocks through neuro-anatomic pathways and diffusible factors involving for example hormones and growth factors. Thus, many cellular functions, including metabolism, cell proliferation and death, display circadian rhythms. Targets of pharmacological agents are also regulated by the CTS. The identification of such targets form the basis of chronopharmacodynamics, which enable, together with chronopharmakokinetics, to determine the optimal time of drug application with the aim to increase efficacy and to decrease adverse effects. This has been especially applied to anticancer drugs. The platinum complex cisplatin for example is best tolerated following dosing near the middle of the active phase in mice and in patients. Nevertheless, cisplatin tumour resistance still remains an issue. To address this problem, a new generation of osmium complexes has been developed and has been proven to overcome platinum resistance in some preclinical models. A main representative of this family is FY26. The research conducted here sought to link in vitro and in vivo chronopharmacology and chronotoxicity of FY26 in preclinical models. For such purpose, the physiological core body temperature rhythm was taken as a circadian biomarker for referring the time points corresponding to best FY26 tolerability.

This research question was addressed first using murine hepatocarcinoma cells (Hepa1-6 Per2-luc) with or without silenced Bmal1, and a human colon cancer cell line (HCT116 Per2-luc). Cell cultures were synchronised with 24-h temperature cycles mimicking the physiological rhythmic variations. Time dependencies were sought for FY26 cytotoxicity, cellular uptake, proliferation, cell cycle phase distribution, apoptosis and autophagy. Towards this goal, FY26 was administered at 4 to 6 circadian time points in synchronised cell populations.
In vitro and in vivo experiments aimed to determine the best tolerated and therapeutic FY26 doses. These doses were further used to assess the circadian tolerability profile and the difference in antitumor efficacy, when given at the respective time point of highest and lowest tolerability. According to the results of the in vitro chronotoxicity study, FY26 was best tolerated at T22, referring to the 2 h before the programmed temperature rise from 36 to 37°C. The in vivo chronotoxicity study identified FY26 to be best tolerated at ZT06. This time point referred to the middle of the inactive phase of mice at which core body temperature was 35.8 ± 0.3°C. In order to determine possible time dependent differences in tumour growth inhibition by FY26, the drug was injected in 5 repetitive i.p. injections every other day at ZT06 or at ZT18. FY26 showed the highest tumour growth inhibition at ZT18, the time point of least tolerability. Those studies were followed by in vitro chronopharmacokinetics and chronopharmacodynamics studies in order to help explain the circadian rhythm in FY26 toxicity.

In vitro pharmacokinetics studies demonstrated maximum and minimum intracellular FY26 uptake at T02 and at T14 respectively. Under continued FY26 exposure for 72 h, the doubling time of Hepa1-6 Per2-luc cells was twice as long following FY26 addition at T18 as compared to that at T10 or T14. Monitoring Per2-luc bioluminescence revealed that the inhibition of Per2-luc bioluminescence was highest following drug addition at T22. Such finding was coherent with the prolonged doubling time resulting from FY26 exposure starting at T18. Flow cytometry analysis revealed an accumulation of FY26-treated cells in S-phase. Such S-phase arrest was larger following FY26 addition at T16 as compared to T04. Taken together the results suggested that FY26 tolerability was best after drug exposure onset at T22, because a higher number of cells transiently accumulated in S-phase, but did not undergo apoptosis. Thus drug addition at T04 resulted in a higher count of apoptotic cells referring to the time point of least FY26 tolerability. FY26 further increased apoptotic cell counts in the shBmal1-silenced clone, accounting for a higher FY26 sensitivity. The measurement of the autophagy gene LC3B confirmed the higher sensitivity of the shBmal1-silenced cell line as compared to Hepa1-6 Per2-luc as shown with higher LC3B mRNA expression levels.
Finally, *in vitro* and *in vivo* experiments were performed in order to determine possible effects of FY26 on the circadian clock itself, and their temporal relations with core body temperature rhythm as a CTS biomarker. Standard statistics, Spectrum resampling, and Cosinor Analyses were applied to the longitudinal time series data in order to determine circadian parameters and to validate intergroup differences. The administration of FY26 at T04, T010, T16 and T22 markedly reduced bioluminescence over the 2.5 to 5 days following drug addition, with significant time-dependency. FY26 did not ablate *Per2-luc* circadian oscillation yet it dampened its amplitude with the highest bioluminescence inhibiting at T22. However, the mRNA expressions of clock genes *Per2*, *Bmal1* and *Rev-erba* were minimally altered by FY26 dosing at T04 or at T22 in synchronised Hepa1-6 *Per2-luc* cells. This suggested that FY26 effects could rather take place at posttranscriptional levels. The *in vivo* relevance of these *in vitro* findings was investigated through the real-time continuous recording of PER2::LUC bioluminescence in transgenic PER2::LUC mice kept in constant darkness. In these conditions, the circadian period of PER2::LUC bioluminescence averaged 22.5 h, so that the real average endogenous circadian times of FY26 dosing were CT24 (range, CT22.5 to CT01.5) and CT12 (CT11.5 to CT13.5). FY26 markedly reduced and altered bioluminescence rhythms for up to 48 hours after injection, with largest amplitude reduction in the mice dosed at CT24. Simultaneously, FY26 altered core body temperature rhythm, an effect which was confirmed in LD12:12 synchronised mice. FY26 injection sharply decreased core body temperature and transiently disrupted its circadian pattern for ~ 24 h following dosing at ZT18. No rhythm alteration was observed at ZT06. The results hence confirm that FY26 can alter the Circadian Timing System, which more susceptible following was dosing at the time of highest systemic toxicity, and least at that of best tolerability.

**In conclusion**, the research presented in this thesis has highlighted the complementary and consistent links between *in vitro* and *in vivo* chronopharmacology findings, as shown for the anticancer drug candidate FY26. The use of 24-hour temperature schedules as an effective synchroniser of circadian clocks *in vitro* and that of core body temperature rhythm as a circadian biomarker *in vivo* has enabled the precise determination of the endogenous circadian times of best tolerability. This was
achieved, as physiological rhythm had been introduced into cell culture conditions. Should these results be applicable to humans, least FY26 toxicity could occur at night, shortly after the physiological nadir in core body temperature. More generally, the synchronisation of healthy and cancer cells with 24-h periodic temperature schedules mimicking physiological cycles could have a high potential to identify optimal drug timing options and underlying chronopharmacokinetics and chronopharmacodynamics mechanisms, whilst reducing and replacing animal experiments, as illustrated here. This could indeed encourage the integration of chronotherapy concepts into drug development strategies. In such context, the objectives of animal chronotherapy studies might hence become mostly confirmatory. Thus, the use of core body temperature circadian cycle appears not only as crucial for the determination of the circadian timing resulting in best drug tolerability, antitumour efficacy, and quality of life, but it is also in agreement with the principles of the 3 R’s (reduction, refinement and replacement) and the public ethical opinion on animal experiments.
### Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AMP</td>
<td>Adenosine monophosphate</td>
</tr>
<tr>
<td>ALT</td>
<td>Alanine Aminotransferase</td>
</tr>
<tr>
<td>APAP</td>
<td>Acetaminophen</td>
</tr>
<tr>
<td>AST</td>
<td>Aspartate Transaminase</td>
</tr>
<tr>
<td>AUC</td>
<td>Area under the curve</td>
</tr>
<tr>
<td>C&lt;sub&gt;max&lt;/sub&gt;</td>
<td>Maximum or peak of concentration</td>
</tr>
<tr>
<td>CREB</td>
<td>Cyclic AMP response element</td>
</tr>
<tr>
<td>CT</td>
<td>Circadian time under constant darkness</td>
</tr>
<tr>
<td>CTS</td>
<td>Circadian timing system</td>
</tr>
<tr>
<td>CYP450</td>
<td>Cytochrome P450</td>
</tr>
<tr>
<td>DD</td>
<td>Darkness-darkness or Constant darkness</td>
</tr>
<tr>
<td>DEPC</td>
<td>Diethyl pyrocarbonate</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s Modified Eagles Medium</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethyl sulfoxide</td>
</tr>
<tr>
<td>Exp</td>
<td>Experiment</td>
</tr>
<tr>
<td>GSH</td>
<td>Reduced Glutathione</td>
</tr>
<tr>
<td>H-NMR</td>
<td>Proton nuclear magnetic resonance</td>
</tr>
<tr>
<td>ICP-MS</td>
<td>Inductively coupled plasma mass spectrometry</td>
</tr>
<tr>
<td>IC&lt;sub&gt;10&lt;/sub&gt;</td>
<td>Inhibitory drug concentration causing 10% cell death</td>
</tr>
<tr>
<td>IC&lt;sub&gt;50&lt;/sub&gt;</td>
<td>Inhibitory drug concentration causing 50% cell death</td>
</tr>
<tr>
<td>i.p.</td>
<td>Intraperitoneal</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>IUPAC</td>
<td>International Union of Pure and Applied Chemistry</td>
</tr>
<tr>
<td>L-BSO</td>
<td>L-buthionine-sulfoximine</td>
</tr>
<tr>
<td>LD 12:12</td>
<td>12 h alternating light-dark cycle</td>
</tr>
<tr>
<td>MDT</td>
<td>Maximum tolerated dose</td>
</tr>
<tr>
<td>MESOR</td>
<td>Midline-estimating statistic of rhythms</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate Buffered Saline</td>
</tr>
<tr>
<td>PC&lt;sub&gt;max&lt;/sub&gt;</td>
<td>Maximum photon counts</td>
</tr>
<tr>
<td>PDT</td>
<td>Population doubling time</td>
</tr>
<tr>
<td>PMT</td>
<td>Photon multiplier tube</td>
</tr>
<tr>
<td>RGC</td>
<td>Retinal ganglion cells</td>
</tr>
<tr>
<td>RGE</td>
<td>Relative gene expression</td>
</tr>
<tr>
<td>RHT</td>
<td>Retinohypothalamic tract</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive oxygen species</td>
</tr>
<tr>
<td>RT-BIO</td>
<td>Real-time Biolumicorder</td>
</tr>
<tr>
<td>s.c.</td>
<td>Subcutaneous</td>
</tr>
<tr>
<td>SCN</td>
<td>Suprachiasmatic nuclei</td>
</tr>
<tr>
<td>SEM</td>
<td>Standard error of the mean</td>
</tr>
<tr>
<td>SRB</td>
<td>Sulforhodamine B</td>
</tr>
<tr>
<td>t&lt;sub&gt;max&lt;/sub&gt;</td>
<td>Time to reach C&lt;sub&gt;max&lt;/sub&gt;</td>
</tr>
<tr>
<td>ZT</td>
<td>Zeitgeber Time (English, synchroniser time)</td>
</tr>
</tbody>
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Research Training

Transferrable Skills Modules at Warwick University

- Writing Focused Scientific Articles and Reports CH934
- Science Communication CH954
- Decision Making and Leadership CH955
- Business, Innovation and Commercialisation for Researchers CH957

Training

- Sleep and Circadian Neuroscience Summer School in Oxford, August 2016
- Workshop XV European Biological Rhythm, August 2017 Amsterdam
- Statistics with R Studio, November-December 2016 University of Warwick

Communication to Congresses

Poster

- XV European Biological Rhythm Society Congress in Amsterdam, August 2017
  Title: “Circadian synchronisation of mouse hepatocarcinoma cells with 24 h temperature cycles uncovers in vitro chronotoxicity of novel anticancer drug”

- Academy of Medical Sciences at Warwick University, April 2017
  Title: ‘Towards determining in vitro chronopharmacology of a new generation of anticancer drugs’

- Sleep and Circadian Neuroscience Summer School in Oxford, August 2016
  Title: ‘Synchronisation properties of temperature cycles in cancerous cell lines’
• GI Circadian Meeting in Chicago, May 2016
  Title: “Toward determining in vitro chronopharmacology of the mTor inhibitor Everolimus in murine hepatocarcinoma cells”

• Poster presentation at the UK Clock Club in Edinburgh, December 2015
  Title: ‘Entrainment of cancer cell lines by temperature cycles’

Talks

• Warwick Medical School PGR Symposium, June 2018
  Title: “It’s all about timing – Impact of the circadian clock on the development of a novel anticancer drug candidate”

• GI Circadian Meeting in Chicago, May 2016
  Title: ‘Toward determining in vitro chronopharmacology of the mTor inhibitor Everolimus in murine hepatocarcinoma cells’

Publications


Teaching

• Facilitating 2nd year Medical School students in case base learning (CBL1, CCE AD and AC1 courses) at Warwick Medical School, October 2017 until now
I Introduction
1 A screenshot of the public perception of circadian rhythm

“Google, what is circadian rhythm?”

“A circadian rhythm is any biological process that displays an endogenous, entrainable oscillation about 24 hours”

This answer is the first definition of circadian rhythm google presents. It was also probably the first answer many people read after researching the meaning of circadian rhythm following the announcement that Jeffrey C. Hall, Michael Rosbash and Michael W. Young were awarded with the Nobel Prize in Physiology or Medicine on the 2nd October 2017.

Figure I-1: Screenshot of the results of the worldwide Google Trend search “circadian rhythm” during the past 12 months (source: Google Trends).

As shown in Figure I-1 the public interest in circadian rhythm was high but short. Nevertheless, this announcement had a huge impact on medical research as it triggered high interest in the possible relevance of circadian rhythm for sleep disorders, and mental health, as well as neurodegenerative, metabolic and malignant diseases. Further, it motivated non-circadian researchers to implement the circadian clock into their field of interest. Unfortunately, less attention has been paid to the impact of the circadian rhythm for pharmacology and drug development. Hence, this thesis can be understood as a contribution to the integration of the circadian rhythm into pharmacology; and in this case into the pharmacology of anticancer agents.
2 Circadian rhythm

The word *circadian* arises from the Latin words *circa* and *diem* and means *approximately a day* (Halberg, 1969). Circadian rhythms are biological variations which recur approximately every 24 h, and whose regulation is reset to 24-h by the regular alternation of natural light and darkness. Rhythms with different oscillatory lengths or periods have been described. For example cyclic adenosine monophosphate (cAMP) is released in a pulsatile manner triggering the release of hormones. Hormones like insulin, gonadotropin releasing hormone (GnRH), growth hormone (GH) are released every 13 min, every hour and every 3 to 5 h. Hence, two kinds of biological rhythms have been distinguished: the pulsatile and the circadian rhythms (Goldbeter, 2002). Regardless of the length of an oscillation, the parameters period, mesor, amplitude and acrophase have been commonly used to describe biological rhythms. The period is the time needed to complete one oscillation. The mesor represents the mean of the fitted cosine function, the amplitude measures the difference between the mesor and the maximum value of the oscillation, and the acrophase is the timing of the maximum. The cosinor and other spectral methods provide these parameters with their confidence limits (Reinberg, 1992; Vitaterna et al., 2001).

In physiology, circadian rhythms can be found in sleep and wake pattern, as well as e.g. body temperature (Dibner et al., 2010). In mammals, physiological circadian rhythms are endogenous, entrainable, self-sustained and temperature compensated. They are moreover generated by circadian clocks (Vitaterna et al., 2001).

3 The circadian timing system

Circadian clocks are conserved amongst different tissues such as liver, kidney, heart, hypothalamus and fibroblasts (Plikus et al., 2015; Zhang et al., 2014). They are organised in a hierarchical network structure resulting in the Circadian Timing System (CTS) (Figure I-2). Exogenous or environmental signals such as light or social cues, serve as *Zeitgeber* (German word for “time giver”) through resetting the central clock in the suprachiasmatic nuclei (SCN). The SCN generates output signals coordinating the circadian clocks in peripheral tissues. Such output signals or internal time cues can
be e.g. hormones and body temperature (Lévi et al., 2010). Melatonin, a sleep hormone, peaks near 04:00 in humans and is involved in the regulation of the sleep-and wake cycle (Cajochen et al., 1999; Cajochen et al., 2003). Corticosterone in rodents and cortisol in humans are rhythmically secreted by the adrenal gland, with peaks near the start of the activity phase of the rest-activity cycle in both species. The glucocorticoid rhythms participate in the circadian regulation of metabolic function and adaptive stress response (Son et al., 2008).

Core body temperature follows a circadian rhythm with highest values at night in rodents and during daytime in humans (Cajochen et al., 2003). Daily changes of core body temperature entrain cell-autonomous peripheral clocks in vivo. The SCN of rodents remains unaffected by external temperature cycles, which however do entrain peripheral clocks in lungs and pituitary tissues, among others. This demonstrates that the rhythm in core body temperature, generated by the SCN, is important to entrain peripheral clocks without affecting the SCN (Buhr et al., 2010). A comparison between the synchronisation properties of light-dark cycles with external temperature cycles in mice, identified temperature cycles as Zeitgeber with weak synchronisation properties (Refindetti, 2010). Nevertheless, temperature cycles mimicking core body temperature were shown to maintain autonomous and self-sustained oscillations in cultured mouse fibroblasts (Brown et al., 2002; Nagoshi et al., 2004; Saini et al., 2012). Several internal time cues provided by the rhythms in core body temperature, circulating melatonin and cortisol levels, are important as they contribute to the coordination of the peripheral clocks machinery. This machinery rhythmically regulates cellular metabolism, proliferation and survival (Dibner et al., 2010; Lévi, 2006).
Figure I-2: Schematic representation of the CTS and its interaction with external time cues and other critical cellular functions (Lévi et al., 2010).

4 Photoperiodic Entrainment of the SCN in Mammals

The translation of the photoperiod in chemical signals is partly mediated by melanopsin located in the retinal ganglion cells (RGC) (Hattar et al., 2002; Ruby et al., 2002). Melanopsin and glutamate are both stored in the pituitary adenylate cyclase-activating polypeptide-containing RGC (Hannibal et al., 2002; Hannibal et al., 2000). Retinal ganglion cells innervate the SCN through the retinohypothalamic tract (RHT) (Berson et al., 2002; Moore et al., 1995). Voltage dependent Ca$^{2+}$ channels are activated by membrane potentials triggering synaptic glutamate release in the RHT. The glutamate further initiates the release of postsynaptic Ca$^{2+}$ and protein kinases A in the neurons of the SCN (Irwin & Allen, 2007). In the presence of calcium, calmodulin and the calcium/modulin kinase 2α is activated leading to the phosphorylation of the cyclin AMP responsive element binding protein (CREB) at Ser133 and Ser142 (Gau et al., 2002). The clock genes *Per1* and *Per2* (further described in the next paragraph), host the cAMP-responsive element (CREs) to which CREB binds promoting the expression of PER1 and PER2 which regulate the transcription of CLOCK/BMAL1 and thus reset the circadian clock (Travnickova-
Bendova et al., 2002). Despite Ca$^{2+}$ regulating the clock function, the protein kinase Ca (PRKCA) interacts with PER2 and hence stabilises PER2. This interplay is dependent on the availability of light (Jakubcakova et al., 2007).

5 The wheels of the circadian clock

The discovery of the biological machinery of mammalian clocks was initiated as clock gene Per was first discovered in 1971 by Konopka and Benzer (Konopka & Benzer, 1971). Thirteen years later, Per was identified to be essential for the entrainment of Drosophila to light-dark cycles (Bargiello et al., 1984; Zehring et al., 1984). Intensive research led to the description of the molecular circadian clock in mammals being constituted with interconnected negative and positive transcription/translation feedback loops (Figure I-3). The molecular clock involves approximately 15 genes. Bmal1 (brain and muscle Arnt-like protein) and Clock (circadian locomotor output cycles kaput) are core genes whose transcriptional activity governs the positive limb of the clock and triggers the transcription of clock genes (e.g., Clock, Bmal1, Per1, Per2, Per3, Cry1, Cry2, Reverbα and Reverbβ). In the nucleus, the bHLH-PAS proteins BMAL1 and CLOCK/NPAS2 heterodimerise. The heterodimers such as CLOCK:BMAL1 bind to E-box elements being short DNA sequences (5'-CACGTG-3'/ 5'-CACGTT-3') in the promoter region of the target gene. Once the CLOCK:BMAL1 heterodimer is linked to the specific sequence, transcription of the targeted clock genes is activated (Gekakis et al., 1998; King et al., 1997). Those clock genes are for instance Per (Period) and Cry (Cryptochrome). They form a negative limb as their proteins, PER and CRY, feedback to Bmal1 and Clock/Npas2 inhibiting their transcriptional activity. Consequently, PER and CRY are responsible for the down regulation of their own transcription (Kume et al., 1999; Sangoram et al., 1998). Per and Cry consist of Perl, Per2, Per3 and Cry1 and Cry2 genes. Their proteins display different repressor activities on the positive limb. It has been shown in Per3 knock out mice, that the rhythmic expressions of Perl, Per2, Cry1 and Bmal1 were similar to those found in Per3 wildtype (Shearman et al., 2000). The study then suggested that Per3 was not essential for maintaining circadian rhythmicity, whereas Perl, Per2, Cry1 and Cry2 were indeed as their silencing disrupted the transcriptional-translational feedback loop and thus circadian rhythmicity (Van Der Horst et al.,
Another feedback loop is generated through the transcription of the clock genes \textit{Rev-erba/β} and \textit{Rora/β,γ}, which are activated by the CLOCK:BMAL1 complex. REV-ERBa/β represses the activation of \textit{Clock} and \textit{Bmal1} presenting a second negative feedback loop. In the meantime, RORα/β competes with REV-ERBa/β targeting the same binding side. Consequently, RORα/β dampens the repressing effect of REV-ERBa/β on \textit{Clock/Npas2} and \textit{Bmal1} transcription (Guillaumond \textit{et al.}, 2005). This interacting dynamic molecular system of activating and inhibiting loops forms the basis of circadian gene expression.

\textbf{Figure 1-3: Molecular makeup of the circadian clock (Dallmann \textit{et al.}, 2014).}

This interplay between negative and positive transcription and translational feedback loop regulates the circadian expression of so-called clock-controlled genes and proteins through transcription factors. Transcription factors DBP, TEF and HFL have a PAR-domain basic leucine zipper (PAR bZip) and regulate the circadian expression of metabolic genes such as \textit{Cyp2a5} and \textit{Cyp2c50}. The P450 oxidoreductase (POR) exhibits a strong circadian rhythm which is suppressed in PAR bZip knock out mice (Gachon \textit{et al.}, 2006). The heat shock factor 1 (HSF1) is a further transcription factor regulating the expression of \textit{HSP1} at the onset of darkness by binding to the heat shock element (HSE) (Reinke \textit{et al.}, 2008). The release of hematopoietic stem cells (HSC) is mediated by the rhythmic release of noradrenaline by the sympathetic nervous
system (SNS) which is governed by the central clock. Noradrenalin down regulates Cxcl12 through the inhibition of the Sp transcription factor (Méndez-Ferrer et al., 2008). The circadian clock further regulates protein expression indirectly through the regulation of protein degradation processes. One example is the rapamycin target mTOR. It is shown that the rhythmic expression of mTOR is regulated by Fbxw7 (F-box and WD-40 domain protein 7) whose expression is further regulated by the clock output gene Dbp. Consequently, FBXW7 was found the exhibit a circadian pattern which are in antiphase to that of the mTOR expression. This suggested that the ubiquitination of mTOR rather than its mRNA expression, was clock-controlled (Okazaki et al., 2014).

6 Implementation of circadian clocks in pharmacology

As demonstrated for mTOR, the circadian clock is involved in the regulation of the circadian expression in many genes and proteins. In this context, it is reported that 43% of the protein-coding genome in mice is rhythmic (Ptitsyn & Gimble, 2011; Zhang et al., 2014). A further study identified that 25% of phosphopeptides in the liver presents with diurnal rhythms (Robles et al., 2017). The identification of the circadian expression of such proteins is crucial as they often represent targets for pharmacological agents, e.g. everolimus targets mTOR, forming the base of chronopharmacology (Okazaki et al., 2014).

Chronopharmacology is the science whose subject it is to determine and describe the time dependent drug effects, taking into account the circadian clocks. This includes studies on circadian changes in pharmacokinetics and pharmacodynamics. Pharmacokinetics describes the properties of drug absorption, distribution, metabolism and elimination (ADME), whereas pharmacodynamics defines the effect of agents with e.g. molecular and cellular targets.

7 Circadian Pharmacokinetics

Gastric pH and motility influence drug absorption thus, determine the speed and quantity of drug uptake. The rhythmicity of gastric pH has been reviewed previously pointing out its importance for drug ionization and hydrophobicity. Additionally, gastric motility is increased during day time as compared to night time in humans.
(Dallmann et al., 2014; Ohdo, 2010). The peptide ghrelin governing gastric motility is regulated by clock gene Bmal1. Ghrelin mRNA expression is highest at circadian time 4 (CT4, beginning of rest span) in wildtype mice kept in DD. This rhythm is ablated in Bmal1 knock out mice (Laermans et al., 2015). A study on 16 healthy male subjects confirmed that the mean emptying half-time is on average 30 min faster in the morning at 08:00 as compared to evening at 20:00 (Goo et al., 1987). The intestinal H+ coupled peptide transporter (PEPT1) mediates the cellular uptake of active drugs such as β-lactam antibiotics or angiotensin-converting enzyme (ACE) inhibitors. The expression of PEPT1 is regulated by clock output gene and transcription factor Dbp with a peak at ZT08 in mice (Saito et al., 2008).

The distribution of pharmacological agents is dependent on blood flow, cardiac output and plasma proteins. Blood flow as well as heart rate exhibit a circadian rhythm with a peak during the late afternoon (Kaneko et al., 1968). The vascular function is clock regulated. A reduced endothelium relaxation and an increase in the expression of cyclooxygenase-1 (COX1) was observed in the aortic ring of Per2 mutant mice but not in wild type mice (Viswambharan et al., 2007). The response of α-adrenoreceptors to phenylephrine and the baroreflex sensitivity are higher during the night in mice. Those rhythms, however, were depleted in Cry1 and Cry2 knockout mice (Masuki et al., 2005). Further, the hepatic blood flow was shown to exhibit a circadian pattern with a peak at 04:00 in the morning (Lemmer & Nold, 1991). The rhythms of hepatic blood flow is influenced by drugs as nifedipine or propranolol as well as food (Feely, 1984; Olanoff et al., 1986). The availability of plasma albumin follows a circadian pattern. The highest plasma concentration of albumin was determined at ZT15 to ZT18 in mice (Mauvoisin et al., 2014) and near 16:00 to 20:00 in human subjects (Jubiz et al., 1972).

Drug metabolism involves phase I and phase II enzymes and phase III transporters. Proteins belonging to phase I enzymes are oxidases, reductases and hydrolases as part of the microsomal cytochrome P450 (CYP450) (Ferrell & Chiang, 2015). They activate or inactivate drugs before undergoing phase II metabolism. In phase II, the hydrophilicity of drugs is increased by their conjugation through UDP-glucuronosyltransferases, N-acetyltransferases, glutathione S-transferases, or thiopurine S-methyltransferases (Jancova et al., 2010). Phase III transporters involve
multiple families, including the ABC transporter P-glycoprotein (P-gp) and multidrug resistance-associated proteins (MRP) as well as the organic anion transporting polypeptide 2 (OATP2) (Xu et al., 2005).

Transcription factors like aryl hydrocarbon receptor (AhR), constitutive androstane receptor (CAR), pregnane X receptor (PXR) or proliferator-activated receptor-α (PPARα) exhibit circadian patterns thus generate and propagate circadian changes in the mRNA expression of Cyp genes. PPARα activates the rhythmic transcription of Cyp4a10 and Cyp4a14. Comparing the transcription levels of Cyp genes in male and female mice liver, revealed a higher expression of Cyp genes in female compared to male mice. Female mice exhibited a 20-fold higher RNA expression of Cyp4a14 as compared to male mice (Lu et al., 2013). In the hepatocellular carcinoma cells, HepG2, the transcription factor Dbp (Yamaguchi et al., 2000) shows a similar circadian oscillating pattern as that in Cyp3A4. Increased DBP levels, mediates the expression of Cyp3A4 in HepG2 cells (Takiguchi et al., 2007). Contrary to Cyp3A4 activation, the circadian expression of the transcription factor PPARα which activates enzymes of the Cyp4 family, is governed by the CLOCK:BMAL1 heterodimer. Circadian rhythms are also found for the transcription of enzymes in the Cyp4 family (Lu et al., 2013; Oishi et al., 2005). As a result, the metabolism of the analgesic and antipyretic drug acetaminophen (APAP) also known as paracetamol, is an example for a drug whose toxicity is affected by the rhythmic expression of metabolising enzymes. Due to higher alanine and aspartate transferase (ALT and AST) and lower glutathione levels at 20:00, an increased concentration of the toxic metabolite N-acetyl-p-benzoquinone occurs in mice synchronised with 12:12 h light-dark cycles with light onset at 07:00 and light off set at 19:00 (Kakan et al., 2011). It is further reported that the APAP toxicity is several fold larger and arrhythmic in hepatic Bmal1 knock out mice. Additionally, the study shows a decrease in Cyp2a1 and Cyp1a2 resulted from reduced transcription activity of P450 cytochrome oxidoreductase (Por) in Bmal1 knock out mice (DeBruyne et al., 2014; Johnson et al., 2014). The mRNA expression of the phase III ABC transporter, P-gp and Mrp followed a circadian expression pattern. Abcb1a (Mdr1a) and Abcb1b (Mdr1b) are highest expressed near ZT16 in mouse liver and near ZT12 in mouse intestine, with large sex-related differences (Alper Okyar1 et al., 2018b). The mRNA of Abcb4 (Mdr3) is highest at
ZT08 in liver and at ZT04 in the intestine (Ando et al., 2005). The multidrug resistance-associated protein 2 (MRP2) and the organic anion transporter 1 and 2 (SLC22A1 and SLC221B) are rhythmically expressed with acrophases at the end of the rest phase in mice (Panda et al., 2002)

Regarding the elimination, studies in wildtype mice demonstrated an oscillating pattern of stool output during the active phase (CT20) and an increased muscle contractile response in colon to acetylcholine at the beginning of the dark phase (CT13). Those effects do not occur in Per1/2 knock out mice (Hoogerwerf et al., 2010). Renal elimination is further circadian regulated (Firsov & Bonny, 2010). It is further reported that the glomerular filtration rate is highest during the late afternoon and lowest during the early morning hours in humans (Koopman et al., 1989).

8 Chronopharmacodynamics of anticancer drugs

A prominent target, especially for anticancer drugs, is the cell cycle which regulates, DNA synthesis, mitotic division, cell growth, apoptosis and autophagy. Schematically, the cell cycle exhibits different cell cycle phases which are called G0/G1, S (DNA replication), G2, and M (mitosis). The transition from one cell cycle phase to the next one is regulated by cyclin-dependent kinases (CDK) and cyclins (Morgan, 1995). CDK1 supports the transition from S-phase to mitosis initiated by cyclin A. CDK2 is involved in the transition from G1- to S-phase (Malumbres & Barbacid, 2009). Cell cycle phase transitions from G1 to S and from G2 to M are controlled by the circadian clock, at least via p21 and wee1. For instance, the protein WEE1 whose gene transcription is activated by CLOCK:BMAL1 heterodimer and suppressed by PER and CRY proteins, is shown to be rhythmic. A rhythmic phosphorylation and activation of CDK1 is promoted resulting in the activation of the G2/M transition (Matsuo et al., 2003). Additionally, clock genes Per1 and Per2 interact with the serine/threonine kinase ATM and the proto oncogene mdm2 controlling DNA repair. Disrupting the rhythmicity in clock gene expression due to environmental changes (e.g., chronic jet lag) down regulate the tumour suppressor gene p53. This results in the activation of cell proliferation and the down regulation of DNA repair mechanism as well as apoptosis (Hunt & Sassone-Corsi, 2007; Lévi, 2006). Recent research further demonstrates that the cell cycle is aligned with the circadian clock in individual
cultured mouse fibroblasts with or without time cues (Bieler et al., 2014; Feillet et al., 2014). This coupling is suggested to be altered in cancerous cell lines (Feillet et al., 2015). It is argued that the fast proliferation of tumour cells is due in part to the uncoupling of the cell cycle from the circadian clock, and that entrainment of cancer cells could be a way to slow down tumour growth (Feillet et al., 2015). It is further proposed that mutations in clock genes could result in the dysregulation of the cell cycles thus promoting cancer cells proliferation and tumour progression (Lévi et al., 2007). Therefore, understanding the link between the circadian clock and the cell cycle in healthy and cancerous cells appears critical for mapping the mechanisms that underlies the chronopharmacology of anticancer agents.

Circadian rhythms in tolerability have now been identified for 50 anticancer agents belonging to eight pharmacology classes (Figure I-4) (Lévi, 2006; Lévi et al., 2010). Such classes are the antimetabolites, topoisomerase inhibitors, intercalating agents, cytokines, mitotic spindle poisons, nitrosoureas, small kinase inhibitors and alkylating agents. Representatives of each class exhibit different circadian times of best drug tolerability. For example the topoisomerase inhibitors irinotecan and etoposide display best tolerability at ZT7 whereas the cytokine interleukine-2 is best tolerated at ZT17 (Lévi et al., 2010). The Lethal Dose (LD) 50 of the antimetabolite 5-fluorouracil (5-FU) in mice is higher following drug injection at 11:00 (LD50: 450-500 mg/kg) as compared to 23:00 (LD50: 250-330 mg/kg) in mice synchronised with lights on from 06:00 to 18:00 (Burns & Beland, 1984). A second in vivo study confirms that 200 mg/kg 5-FU exhibit the highest toxicity at 04:00. Thus, 5-FU is best tolerated when administered during the early rest phase of mice (Popovic et al., 1982). The mitotic inhibitor docetaxel is best tolerated in mice following the i.v. injection of 30 mg/kg docetaxel at ZT7 to ZT11. Strikingly, optimal antitumour efficacy of docetaxel is also achieved following dosing near the time of best tolerability. Thus, the inhibition of tumour proliferation is shown to be 2 to 3 times larger following docetaxel injection to pancreatic cancer-bearing male B6D2F1 mice at ZT7 as compared to ZT19 or ZT23 (Tampellini et al., 1998). The time point of best docetaxel tolerability is further confirmed following the injection of 38.8 or 23.3 mg/kg/injection to male C3H/HeN mice. The same study demonstrates that male C3H/HeN mice injected with 13.8 and 8.3 mg/kg/injection of the DNA intercalator doxorubicin lose significantly less body
weight following treatment at ZT7, thus confirming earlier findings of best doxorubicin tolerability in male CDF1 mice (Granda et al., 2001; Sothern et al., 1988). The administration of docetaxel and doxorubicin at ZT7 prolonged tumour growth inhibition by 50% as compared to other circadian time points in breast cancer-bearing C3H/HeN mice. The co-administration of 16.3 mg/kg/injection docetaxel and 2.5 mg/kg/injection doxorubicin once per week for 3 weeks leads to a persistent tumour inhibition of 45% with the best tolerability at ZT7 (Granda et al., 2001). Alkylating agents are another class to display marked circadian toxicity profiles. This is especially the case for platinum complexes cisplatin, carboplatin and oxaliplatin.

Figure I-4: Overview of best timing of anticancer agents.

The figure describes the circadian timing of anticancer agents in 8 pharmacologic classes. Platinum complexes such as cisplatin, carboplatin and oxaliplatin are best tolerated when given during the early hours of the active phase in nocturnal rodents, despite large differences in pharmacology and action mechanisms. The red polar curve depicts the average circadian rhythm in core body temperature (Lévi et al., 2010).

9 Chronopharmacology of platinum complexes

The anticancer activity of platinum compounds was first recognised in 1965 by Rosenberg. He noticed that the bacterium Escherichia coli would stop dividing during
the presence of the cisplatin, that was freed from the platinum electrodes as a result of electric current flow (Rosenberg et al., 1965). Following his discovery, clinical trials identified cisplatin as an anticancer drug which received FDA approval in 1978 (Monneret, 2011). Platinum compounds like cisplatin, carboplatin and oxaliplatin today represent the most widely used drugs to treat cancer. Cisplatin is approved for the treatment of ovarian, breast, bladder, lung, head and neck carcinoma in combination with a taxane. It is further used for the treatment of gastric and esophagogastric adenocarcinoma in combination with paclitaxel and 5-FU as well as against prostate, cervical and biliary cancer (Dasari & Tchounwou, 2014). However, severe adverse effect such as renal failure, anemia, neutropenia and nausea are limiting the use of cisplatin. Thus, the determination of the optimal cisplatin timing could contribute to the improvement of its tolerability.

Initial studies in rats demonstrate a several fold improvement in cisplatin gross, renal and haematological toxicities through the delivery of this drug near ZT18, i.e. near the middle of the nocturnal active span in female F344 Fisher rats (Hrushesky et al., 1982b; Levi et al., 1982). Interestingly the antitumour efficacy of cisplatin (in combination with doxorubicin) is highest following dosing near the middle of the nocturnal activity span in tumour-bearing rats (Soehn et al. 1989). Similar studies investigated the chronotolerance of cisplatin in male B6D2F1 mice and revealed that the time point of best drug tolerability is still located during darkness, yet slightly earlier, i.e. near ZT15-16 (Boughattas et al., 1989). Subsequently, an experiment determining the tolerability and the antitumour efficacy of gemcitabine (deoxycytidine analogue) in combination with cisplatin in tumour-bearing mice is performed. ZT15 and ZT11 are identified as the most effective dosing times of cisplatin and gemcitabine respectively, both as single agents and in combination (Li et al., 2005b).

Subsequent chronopharmacology studies demonstrate similar chronotolerance and chronoefficacy patterns for carboplatin and oxaliplatin, with a similar “optimal” timing at ZT15-ZT16 in male B6D2F1 mice, despite major differences in toxicity target tissues, pharmacokinetics, and pharmacodynamics (Boughattas et al 1988, 1989, 1990). While chronopharmacokinetics do not appear to play a major role, the circadian detoxification of these compounds with reduced glutathione (GSH) is critical. The inhibition of glutathione synthesis by the glutathione synthesis inhibitor
buthionine sulfoximine (BSO) increases jejunal toxicity and oxaliplatin intestinal uptake (Lévi et al., 2000). Another study searched for the drug administration time associated with best antitumour efficacy of oxaliplatin, using Glasgow osteosarcoma-bearing mice. A daily dose of 5.25 mg/kg of oxaliplatin achieves both best tolerability and best efficacy at ZT15 compared to ZT7 (Granda et al., 2002). In aggregate the delivery of cisplatin or oxaliplatin near the middle of the nocturnal active span of rats or mice demonstrate both best tolerability and best antitumour efficacy. However, sex and possibly species or strain could moderate their chronopharmacology to some extent.

Clinical chronotherapy trials investigated the clinical relevance of these findings, with regard to both tolerability and efficacy in cancer patients. Initial randomised trials have shown that cisplatin is less toxic and possibly more effective following dosing in the late afternoon hours (16:00-18:00) as compared to early morning hours (04:00-06:00) in ovarian cancer patients (Hrushesky et al., 1982a; Levi et al., 1990). Subsequent clinical trials mostly investigate the relevance of oxaliplatin timing, in combination with the anticancer drug 5-fluorouracil (5-FU) and leucovorin in patients with metastatic colorectal cancer. The chronotherapy trials were the first ones to demonstrate any clinical efficacy of a platinum complex in human colorectal cancer (Levi et al. 1992). In two international randomised trials, oxaliplatin chronotherapy proved as less neurotoxic and more effective as compared to flat infusion (Levi et al. 2001). Subsequently, a combination of 700 - 1100 mg/m²/day 5-FU (peak at 04:00), 300 mg/m²/day leucovorin (peak at 04:00) and 25 mg/m²/day oxaliplatin (peak at 16:00) is infused as circadian chronomodulated infusions for up to 12 courses in patients with metastatic colorectal cancer. The chronomodulated dosing schedule allows an increase in dose intensity for 5-FU up to 36%. Moreover the same schedule improves objective response rate up to 15% suggesting that chronotherapy has an impact on minimising side effects and improving the outcome of pharmacotherapy (Lévi et al., 1999). This study was followed by a clinical phase III trial comparing chronomodulated delivery to a non-time stipulated conventional drug administration of the same drugs in 554 patients (338 male and 226 female). Safety, response and survival are monitored. The results report no difference in overall survival between the chronotherapy schedule and the conventional schedule. Yet, chronotherapy
improved haematological tolerability several fold and it significantly prolonged survival in men, but not in women as compared to conventional delivery (Giacchetti et al., 2006).

The sex-chronotherapy schedule interaction is further confirmed in a meta-analysis of three randomised international trials comparing chronotherapy vs. conventional delivery (Giacchetti et al. 2012). Additional experimental studies and clinical trials suggest that the optimal timing could differ by nearly 6 h between males and females (Levi et al. 2007; Li et al. 2013). Ongoing research is aiming at the personalisation of chronotherapy delivery according to both sex and the CTS of cancer patients, using tele-monitored circadian biomarkers and a systems chronopharmacology approach (Ballesta et al, Pharm Rev 2017). Further, in silico models paired with in vitro models are promising new methods to assess the best time of drug administration through a system approach. This was first developed for irinotecan chronopharmacology. By applying a comprehensive mechanistic circadian PK-PD model on chronopharmacokinetic and chronopharmakodynamic data on irinotecan, the critical circadian rhythms in proteins involved in the mechanism of action are identified (Dulong et al., 2015).

Another variable affecting the determination of the optimal timing of cancer drugs is the disturbance of the circadian clock (Ballesta et al., 2011). CTS disruption is associated with a lower survival rate in cancer patients. For example the disruption of daily cortisol patterns is related to a higher risk of an earlier mortality in patients with metastatic breast cancer (Sephton et al., 2000). A poorer survival is also demonstrated in patients with metastatic gastro-intestinal malignancies and circadian disruption of their rest-activity or body temperature rhythms (Ballesta et al., 2017). On the other hand, anticancer drugs could disrupt the CTS through altering core body temperature and/or rest-activity rhythms (Ballesta et al., 2017; Lévi et al., 2010). Hence, the determination of optimal drug timing should encompass both the identification of the least toxic drug administration time, but also the quantification and dosing time dependency in the effect of the anticancer drug on the circadian clocks. Other factors such as sex, age and feeding should also be considered jointly, as they could impact on the translation of the basic chronopharmacology findings toward their clinical applications (Ballesta et al., 2011; Innominato et al., 2010).
10 Organometallic osmium compounds FY25 and FY26

In order to attempt at improving the therapeutic outcome and survival of cancer patients, a new generation of organometallic anticancer compounds entered the pipeline of drug development. These complexes involve different metal ions, including ruthenium, iridium and osmium (Fu et al., 2010; Liu et al., 2011; Novakova et al., 2003; Novakova et al., 2005). Iridium complexes have been the first compounds to demonstrate an anticancer activity different from platinum and ruthenium complexes. The findings suggested that the main mechanism of action is based on the induction of high ROS levels leading to the activation of cell death mechanisms through calcium release from the endoplasmic reticulum, which further destabilises mitochondria and initiates cell death mechanism (Liu et al., 2014; Liu & Sadler, 2014; Sanchez-Cano et al., 2017).

Table I-1: IUPAC name of the organo-osmium compounds FY25 and FY26 (van Rijt et al., 2014).

<table>
<thead>
<tr>
<th>Compound</th>
<th>IUPAC name</th>
<th>Arene</th>
<th>R</th>
<th>X</th>
</tr>
</thead>
<tbody>
<tr>
<td>FY25</td>
<td>[Os(η⁶-p-cym)(phenylazopyridine-NMe₂)Cl]PF₆</td>
<td>p-cymene</td>
<td>H</td>
<td>Cl</td>
</tr>
<tr>
<td>FY26</td>
<td>[Os(η⁶-p-cym)(phenylazopyridine-NMe₂)I]PF₆</td>
<td>p-cymene</td>
<td>H</td>
<td>I</td>
</tr>
</tbody>
</table>

The main representatives of the organo-osmium compounds are FY25 and FY26. Both compounds display an aromatic hydrocarbon, also called arene, which binds to osmium as a mutual skeleton. FY25 and FY26 consist of the ligands p-cymene and phenylazopyridine bound to osmium. FY25 and FY26 can be distinguished through their fourth ligand which is the chloride for FY25 and the iodide for FY26 resulting in different chemical and biological behaviour (Figure I-5 and Table I-1).
Regarding their chemical properties, both compounds are soluble in water-based culture medium over a period of 24 h. Further, 50 µM of both compounds were diluted in a 10 mM phosphate buffer with deuterium water and then screened for structural changes with 1H NMR. After an incubation time of 24 h, the H NMR spectra showed no structural changes demonstrating a stable molecular structure of both compounds. Results of the ESI-MS analysis indicate a strong osmium chloride bound as no hydrolysis in a 10% methanol/90% water solution occurred (Fu et al., 2010). Due to their stability, FY25 and FY26 were used for biological activity tests. The IC<sub>50</sub> of FY25 and that of FY26 were by 2- to 3-fold and 10- to 15-fold lower, respectively, as compared to the IC<sub>50</sub> of cisplatin in human lung cancer cells (A549), in human colon cancer cells (HCT116) and in human breast cancer cells (MCF-7) (Fu et al., 2010). Additionally, FY26 exhibits improved selectivity towards cancerous cells as compared to cisplatin. The IC<sub>50</sub> ratio between fibroblasts (MRC5) and ovarian cancer cells (A2780) is reported to be 28 for FY26, as compared to 9.5 for cisplatin, thus demonstrating higher selectivity and enhanced anti-proliferative effects (Romero-Canelón et al., 2015). To address the recurring problem of resistance after cisplatin treatment, the IC<sub>50</sub> of FY25, FY26 and cisplatin were compared using cisplatin resistant and cisplatin sensitive human ovarian cancer cells (A2780-cis and A2780, respectively). In sensitive A2780 cells, the IC<sub>50</sub> of FY26 (0.18 ± 0.01 µM) is 10-fold as low those of FY25 or cisplatin (1.8 ± 0.1 µM). Interestingly, in the cisplatin-resistant ovarian cancer cells, the IC<sub>50</sub> of FY25 (1.77 ± 0.68 µM) and that of FY26 (0.23 ± 0.05 µM) do not increase, whereas that of cisplatin doubles (4.03 ± 2.08 µM).
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(Fu et al., 2010). This finding indicate that the cytotoxicity of both Osmium complexes is not altered by cisplatin resistance.

FY25 and FY26 are rapidly taken up during the first 30 min after administration with maximum intracellular uptake after 24 h in cultured human lung cancer cells (A549). Despite this observation, the measurement of the intracellular osmium concentration reveals a 3-fold larger uptake of FY26 as compared to FY25. This phenomenon could be explained with results from another study detecting free extracellular iodide that is released from FY26. However, no free extra- or intracellular chloride is released from FY25 (van Rijt et al., 2014). Intracellular FY26 is surprisingly bio-activated to FY26-OH through hydrolysis, which is mediated by the antioxidant glutathione (GSH) (Fu et al., 2010; Needham et al., 2017). The inertness of the osmium-iodide bound in FY26 explains the high intracellular uptake as well as the amplified biological activity as compared to FY25. Due to the higher reactivity of the osmium-chloride bound, FY25 already starts to undergo hydrolysis before the start of its cellular uptake, which reduces the intracellular amount of FY25. The hydrolysis might not necessarily be mediated by GSH but might be due to the higher reactivity of the chloride bond (Needham et al., 2017). GSH is known to act as an intracellular antioxidant which compensates and protects the cell against reactive oxygen species (ROS) and binds to and detoxifies metallodrugs like platinum compounds (Lu, 2013; Needham et al., 2017). A reduction in GSH levels through its reaction with hydrolysed FY26 weakens cellular defence against ROS, thus increasing ROS production after FY26 administration as shown in cultured A2780 cells. Exposing them to the IC50 of FY26 (0.16 µM) for 24 h increases ROS levels (Hearn et al., 2015). ROS levels in fibroblast (MRC5) only increased following the exposure to 28 time the FY26 IC50 of A2780 cells (Romero-Canelón et al., 2015). This might be due to the intact mitochondrial activity in the fibroblasts which could balance the redox state, whereas the mitochondria in cancerous cells displays altered mitochondrial function (Gogvadze et al., 2008). GSH counterbalances excessive ROS production, thus restoring homeostasis (Armstrong et al., 2002; Mailloux et al., 2013). The co-administration of 5 µM L-buthionine sulfoximine (L-BSO), a redox modulator which inhibits GSH biosynthesis, further elevates ROS production in A2780 cells, and lowers the IC50 of FY26 from 0.16 µM to 0.069 µM ± 0.005 µM. This effect is reversed by addition of
GSH, thus emphasising its crucial role as a potential target of FY26. High levels of ROS are further observed under the co-administration of L-BSO (Figure I-6) (Romero-Canelón et al., 2015).

![Figure I-6](image)

**Figure I-6: Selectivity of FY26 towards non-cancerous cells and ROS induction.**

(A) The co-administration of L-BSO is further able to minimise the IC50 of FY26 and thus increase the selectivity towards non-cancerous cells. (B) FY26 induces ROS production at IC50 concentrations. The co-administration of L-BSO further elevates ROS levels. Compared to non-cancerous cells, FY26 lead to a minor increase in ROS levels (Romero-Canelón et al., 2015).

Cellular distribution experiments reported an increased accumulation of FY26 in cellular membranes, including the mitochondrial ones, 24 h after drug exposure onset. About 74-80% of osmium cellular content is detected in mitochondria membranes, whereas only 4-15% of total osmium content is localised in the cytosol, nucleus and cytoskeleton (van Rijt et al., 2014). Additionally, a high extracellular calcium concentration is triggered with FY26 exposure. Nucleus defragmentation and membrane blebbing are further observed. It is suggested that in response to the high ROS levels resulting from FY26 treatment, calcium is released from the endoplasmic
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reticulum contributing to mitochondrial dysfunction and the activation of cell death mechanism (Figure I-7) (Sanchez-Cano et al., 2017).

Figure I-7: Hypothesised mechanism of action of FY26.

After being taken up into the cell, FY26 undergoes hydrolysis mediated by GSH. Thus bioactive FY26-OH as well as thiolato (GS-) and sulfenato (GSO-) adducts are formed. GSH, as an antioxidant, balanced ROS levels. Depleted ROS levels leads to further production of ROS, which causes calcium release from the endoplasmic reticulum. High cytosolic calcium levels contribute to mitochondrial dysfunction and promote the initiation of cell death mechanisms (Sanchez-Cano et al., 2017).

The ability of FY26 to induce apoptosis is assessed in A2780 (ovarian cancer) and A549 (lung cancer) cells by exposing them for 24 h to their respective FY26 IC50 concentrations. No increase in the population of apoptotic cells is observed after this timespan. The exposure of A549 cells to FY26 concentrations that are 10 times higher than the IC50 only merely increased the number of apoptotic cells. Co-administration with L-BSO results in higher rates of apoptotic cell deaths (Romero-Canelón et al., 2015; van Rijt et al., 2014). However, exposure of A549 cells to 1 µM of FY26, activates cytochrome c release, a marker of early apoptosis, without cells completing apoptosis. Cell cycle analysis reveals an arrest in G1-phase after exposing A2780 cells to 0.16 µM (IC50) to. The same observation is found in A549 cells exposed to 2.5 µM. After the exposure to 5 µM FY26 an accumulation of cells in S-phase is observed,
which is also reported for platinum complexes (Romero-Canelón et al., 2015; van Rijt et al., 2014). In vivo studies, compared the efficacy of FY26 to that of cisplatin. Colon cancer (HCT116) xenografted mice received a single injection of 40 mg/kg of FY26 or 8 mg/kg of cisplatin (Shnyder et al., 2011). Tumour growth was determined for 13 days after drug administration. A significant delay in tumour growth is demonstrated in FY26-treated mice as compared to both untreated controls and cisplatin-treated animals (Shnyder et al., 2011).

![Figure 1-8: Tumour growth inhibition of human colorectal cancer cell line (HCT116) in nude mice.](image)

Forty mg/kg of FY26 and 8 mg/kg of cisplatin were injected as a single dose in tumour bearing mice. Tumour growth was monitored over 13 days revealing a significant delay in the tumour growth after FY26 administration as compared to cisplatin and the untreated control (Shnyder et al., 2011).

The pharmacokinetics and pharmacodynamics investigations suggest a different mechanism of action of the organo-osmium compound FY26 as compared to platinum complexes. Although FY26 causes an increase of S-phase cells, the main mechanism of action of FY26 was different from that of the platinum drugs. Whilst GSH binding to Pt complexes detoxifies them, GSH binding to FY26 forms the bioactive adduct
FY26-OH which increases intracellular levels of ROS. ROS production triggers calcium release from endoplasmic reticulum, which further contributes to mitochondrial dysfunction and initiation of cell death mechanism. However, the mitochondrial target of FY26 still remains to be determined.
11 Scope of the thesis

The aim of my doctoral project is to unravel the chronopharmacology of the potential osmium anticancer drug FY26, so as to optimise tolerability through drug timing and to determine its links to the physiological core body temperature cycles. The chronopharmacological investigations were conducted using *in vitro* and *in vivo* models.

The use of the *in vitro* cell culture models addressed the following research questions:

1. Is temperature synchronisation of cell cultures a suitable method for chronopharmacology studies? And if so, in which conditions?
2. Does FY26 circadian timing *in vitro* alter the extent of its toxicity?
3. If FY26 exhibits a circadian *in vitro* toxicity, what are the main determining factors responsible for its *in vitro* chronotoxicity?
4. Does FY26 affect the circadian clock?

The studies that were performed in mice sought to answer the questions:

5. Is there a consistent circadian rhythm in FY26 tolerability? If so, when is this osmium complex best tolerated?
6. Does FY26 inhibit tumour growth in hepatocarcinoma-bearing mice?
7. Does FY26 antitumour efficacy differ following dosing at the time points of highest or worst tolerability?
8. Does FY26 affect core body temperature and liver PER2 protein expression rhythms when dosed at the time points of best and worst tolerability?
9. Can programmed *in vitro* 24-h periodic temperature cycles be used to predict *in vivo* chronotoxicity, using FY26 as an example?

Answering to those questions is a crucial part not only for the optimisation of FY26 through chronotherapeutic delivery, but also for developmental cancer chronotherapeutics in general. Thus, my research was integrated within a multidisciplinary project that has been seeking to further improve chronotherapy with metallo-drugs through automatic nanoparticle chronodelivery system sensing the circadian rhythm in each cell. This project has involved the research groups of Professor Peter Sadler, Professor Sebastien Perrier, Professor David Rand and Doctor
Annabelle Ballesta. Indeed, the determination of the best time of drug administration constitutes an important aspect for developmental cancer chronotherapeutics as efficacy and survival rates can be improved. Moreover, dosing at the time of highest tolerability increases side effects and thus can improve life quality of patients undergoing chemotherapy.
II Methods and Materials
1 In Vitro Methodology

1.1 Cell lines and cell culture

The *in vitro* chronopharmacology of a new generation of osmium metal compounds was assessed by using an epithelial hepatocarcinoma cell line (Hepa1-6, ATCC® CRL-1830™) arising from BW7756 tumour in C57L mouse and the epithelial colorectal carcinoma cell line (HCT116, ATCC® CRL-247™) from a male adult patient. Modified clones of the Hepa1-6 cell line were engineered within the C5SYS ERASYSBIO+ project (FP7) and were kindly provided, together with the HCT116 cell line, by Dr. Sandrine Dulong from INSERM U935 (Villejuif, France). Hepa1-6 and HCT116 were transfected with the *Per2*-*luc* and *Bmal1*-*luc* reporter construct (Figure II-1). The transfection of the *Bmal1*-*luc* reporter into the HCT116 cell lines was done by our laboratory team within our facility. The gene *Bmal1* was silenced with *shBmal1* RNA (expression of *shBmal1* shown in Chapter III) in Hepa1-6 *Per2*-*luc* by Professor G.T.J. van der Horst (Erasmus University, Rotterdam).

![Figure II-1: Plasmid map of Per2-luc reporter construct inserted in Hepa1-6 cells.](image)

*The plasmid contained the sequence of the Per2 promoter (light blue), of luciferase as well as the CMV promoter, 5’ and 3’ LTR, Gag and the Rev Response Element (RRE).*
All cell lines were grown at 37°C, 5% CO₂ and 90% humidity in a cell culture incubator (Memmert INCO 153, Memmert, Schwabach, Germany) or stored at -196°C. Cells were cultured in Dulbecco’s Modified Eagle’s Medium (DMEM) containing 10% heat inactivated fetal calf serum (FCS, Labtech, Heathfield, UK), 100 µg/mL Penicillin/Streptomycin (Prep room, Life Sciences, University of Warwick) and grown in 10 cm Petri dishes (Sarstedt, Nümbrecht, Germany). The media was prepared by the prep room in the Life Sciences Department of Warwick University.

1.2 Cell thawing and freezing

Cells were plated in a 10 cm Petri dish and frozen until further use once reaching 80% confluency. The media was removed by suction. Cells were washed with 10 mL Phosphate Buffered Saline (PBS, Prep room, Life Sciences, University of Warwick) and incubated with 1 mL Trypsin-EDTA (Prep room, Life Sciences, University of Warwick). The incubation time varied from 1 to 5 min in between cell lines. Nine mL media was the added. The 10 mL cell suspension was then transferred into a 15 mL FALCON tube (Sarstedt, Nümbrecht, Germany) and centrifuged at 200 rcf (Relative Centrifugal Field) for 4 min. Following this, the supernatant was removed and the cell pellet was resuspended in 4 mL media [DMEM with 10% DMSO (Mediatech, Inc., Corning, New York, USA)]. Vials were labelled and frozen at -80°C for 24 h before being stored in liquid nitrogen.

To defrost a cell line, a frozen vial was taken out from the liquid nitrogen tank, put on dry ice and transported to the laboratory. The vial with frozen cells was then defrosted in the Stuart water bath (Cole-Parmer, St. Neots, UK) at 37°C for 3 min. The 1 mL cell suspension was transferred into a 15 mL Falcon™ tube. Using a 10 mL serological Fisherbrand™ pipette (Thermo Fisher Scientific, Waltham, USA), 9 mL cell culture media was added gradually to avoid cell death due to high osmotic pressure caused by DMSO. The cells were separated from DMSO containing media by centrifugation for 3 min at 1000 rcf. The supernatant was removed and the cell pellet suspended in 1 mL fresh media. The new cell suspension was pipetted in a 10 cm Petri dish containing 9 mL media.
1.3 Cell counting

Cells were counted with the Countess automated cell counter (Invitrogen, Thermo Fisher Scientific, Waltham, USA). The media was removed from a 10 cm Petri dish and the adherent cells were washed with 10 mL PBS and incubated with 1 mL Trypsin-EDTA. The incubation times ranged from 1 to 5 min according to cell line. Nine mL fresh media was added into the 10 cm Petri dish to stop trypsination and to homogenise the cells. Ten μL of the cell suspension was used to determine the number of cells per millilitre by homogenising it with 10 μL of the diazo dye 0.4% Trypan-Blue (Invitrogen, Thermo Fisher Scientific, Waltham, USA) solution. Half of the volume was pipetted into a Countess counting chamber slide (Invitrogen, Thermo Fisher Scientific, Waltham, USA) and put into the counting chamber. Due to its ability to permeate damaged cell membranes of dead cells, the dye Trypan-Blue is able to exclude living cells as it is not able to permeate the cell membrane (Strober, 2001).

1.4 Generation of reporter cell lines

1.4.1 Lentivirus production

The *Bmal1*-luc (pLV7-Bsd-P(Bmal1)-dLuc) and *Per2*-luc (pLV7-Bsd-P(Per2)-dLuc) reporter construct were introduced in the human colon cancer cell line HCT116 (ATCC® CRL-247™) (Liu et al., 2008; Ramanathan et al., 2012). Highly proliferating HEK293FT cells were cultured in 10 cm Petri dishes in DMEM media containing 10% FCS and 100 μg/mL Penicillin/Streptomycin. A confluent dish of HEK293TF cell was split 1:1 24 h before transfection to ensure 60% to 80% confluency at time of transfection. On transfection day, the Transfection (TF)-DNA complex, was calculated. For a 10 cm Petri dish, media was added to the TF-DNA complex to obtain a final volume of 1 mL which was added later on the HEK293TF cells. The TF-DNA complex was composed of plasmid, 1.918 μg/μL packaging (pCMVR8.74) (Addgene, Cambridge, USA) and 2.496 μg/μL envelop (pMD2.G) (Addgene, Cambridge, USA). The ratio between the three components was 3:2:1. The ration between the TransIT®-Lenti (Mirus BIO LLC, Madison, USA) transfection reagent and the TF-DNA complex was 3:1. A final TF-DNA complex contained thus 7 μg plasmid, 3.5 μg packaging, 1.75 μg envelop and 21 μg TransIT®-Lenti transfection reagent. According to the DNA concentrations, the volumes of the
packaging, envelop and TransIT®-Lenti transfection reagent were calculated and FCS and antibiotic free DMEM media were added to obtain a final volume of 1 mL. The TF-DNA complex was mixed and incubated at room temperature for 20 min. Meanwhile, the media from the HEK293TF cells was replaced with an antibiotics-containing media, to avoid any contaminations. The TF-DNA mix was added, then followed with a 48 h incubation time at 37°C and 5% CO₂. Once the 48 h incubation time was completed, the media containing the lentivirus was collected and centrifuged at 400 rcf for 10 min at 4°C. The supernatant was filtered through a 0.22 µm filter and aliquots of 1 to 5 mL were made and frozen at -80°C for further use.

1.4.2 Lentiviral transduction

With the lentiviral transduction, the Bmal1-luc reporter was introduced into HCT116 cells. HCT116 cells were cultured to reach a confluence of 50% in a 10 cm Petri dish on the transduction day. The media was removed and 1 mL of the previously produced lentivirus solution, with 8 µg/mL polybrene, was pipetted onto the cells and incubated for 6 h at 37°C and 5% CO₂. After the incubation time was completed, the access to the lentiviral solution was replaced by 10 mL DMEM media with 10% FCS and 100 µg/mL Penicillin/Streptomycin. After another 3 to 4 days, the successfully transduced cells were selected from the non-transfected cells using the antibiotic blasticidin (Corning, New York, USA). As the plasmid carried a blasticidin resistance gene, no cell death should occur following the exposure of blasticidin concentrations. Prior to the transduction, HCT116 cells have been subjected to 1 µg/mL, 2 µg/mL, 4 µg/mL, 8 µg/mL, 16 µg/mL or 32 µg/mL blasticidin, followed by a visual examination of living HCT116 cells under the microscope. It was observed that under 8 µg/mL blasticidin cell death occurred in 80 to 90% of the cells (data not shown). No cell survival was seen under 16 µg/mL blasticidin. Thus, a final concentration of 10 µg/mL blasticidin was used for selection. The transduced HCT116 cells were kept under antibiotic selection for 2 weeks before switching to blasticidin free DMEM media.
1.5 In Vitro synchronisation protocols

1.5.1 Defining target cellular conditions for synchronisation protocols

The purpose of these initial experiments was to determine the number of cells as well as the luciferin concentration needed to synchronise a cell population which remained synchronised for at least 6 days. This allowed to conduct circadian experiments, while keeping the cells synchronised to stable periods, acrophases and amplitudes. All parameters were tested by seeding a given cell suspension in 35 mm NUNC Petri dish (Scientific Laboratories Supplies, Nottingham, UK). Such a cell suspension contained 30 000, 50 000, 75 000, 100 000 or 200 000 cells in 3 mL media. These conditions were applied to cell populations which were synchronised either with dexamethasone (see chapter II.1.5.2) or with temperature cycles (see chapter II.1.5.3). Next, the different luciferin concentrations were tested in order to optimise bioluminescence detection. The aim was to define a balanced luciferin concentration which would provide luciferin for a 7 day long experiment and ensure stable periods, acrophases and amplitudes of the synchronised cell population (Feeney et al., 2016). The optimisation of the luciferin concentration was done in 35 mm NUNC Petri dishes seeded with 50 000 Hepa1-6 Per2-luc cells in 3 mL phenol red free media. Upon cell thawing, luciferin was added in the media so as to obtain concentrations of 50 µM, 100 µM, 200 µM, 400 µM, 800 µM and 1.6 mM. Each concentration was tested in three individual dishes. Immediately after seeding, cells in each Petri dish were subjected to temperature cycle D (Figure II-2) as described in 1.5.3, and put into the LumiCycle (Actimetrics, Willmette, USA) to record bioluminescence for one min every 10 min over a period of 6 to 7 days. The data were recorded and analysed as described in the paragraph “Statistical Methods”.

1.5.2 Synchronisation with dexamethasone

Cells were synchronised with a single dexamethasone shock (Balsalobre et al., 2000). Seventy-five-thousand cells in a 3 mL media cell suspension were seeded in a 35 mm NUNC Petri dish. Synchronisation was started by adding 3 µL of a 1 mM dexamethasone to generate a working concentration of 1 nM dexamethasone (Sigma Aldrich, Gillingham, UK). Cells were exposed for 30 min to the dexamethasone-media solution. The Media was then removed by suction. The cells were washed with 1 mL
PBS, then 3 mL phenol red free DMEM (Sigma Aldrich, Gillingham, UK) containing 100 µM luciferin (Promega, Madison, USA) was added to the dish. The 35 mm NUNC Petri dishes were sealed with a thin layer of silicone grease (RS Components Ltd., Corby, Northants, UK) and put into a LumiCycle (Actimetrics, Wilmette, USA) for bioluminescence monitoring (Chapter II.1.7).

1.5.3 Synchronisation with temperature cycles

It has been shown that temperature cycles mimicking the *in vivo* core body temperature rhythms were able to synchronise the circadian clocks of cells in culture (Brown *et al.*, 2002). To further study the synchronisation properties of exogenous temperature cycles in our model systems, a 50 000 to 100 000 cells were seeded in 3 mL phenol red free DMEM with 100 µM luciferin in 35 mm NUNC Petri dishes. Dishes were sealed with silicone grease and put into the LumiCycle. The optimal temperature cycle protocol was determined by testing several different temperature cycles. The temperature cycles differed by (i) period length (*T* = 24 h for schedules A-D, G, H; *T* = 20 h for schedule E; *T* = 26 h for schedule F) and (ii) subsequent entrainment using temperature cycles consisting of 12-h temperature plateaus differing by 1°C (schedules C, D, E and F) or subsequent “free-running” condition at constant 37°C (schedules A and B) (Figure II-2).
Temperature cycles A to D, G and H were programmed to entrain cells to a 24 h period whereas cells subjected to the temperature cycles E and F were entrained to a 20 h and 26 h period respectively. Temperature cycle A (5-5-constant) started at 37°C for 12 h and decreased to 32°C for another 12 h. This cycle was repeated once, then the temperature was kept constant at 37°C. Temperature cycle B (5-constant) started with a temperature plateau at 37°C for 12 h followed by 32°C for another 12 h, then the temperature was kept constant at 37°C. Temperature cycles C (5-5-1-1-1) and D (5-1-1-1-1) involved initial “temperature shocking cycles” similar to those in schedules A and B, followed with 24 h temperature cycles with alternating 12-h-plateaus at 37°C and 36°C. The temperature profiles of schedules E (5-
Methods and Materials

1-1-1-1) and F (5-1-1-1-1) were similar to schedule D except the cycle period which was set at 20 h for schedule E (alternating 10-h temperature plateaus at 37°C and 36°C), and at 26 h for schedule F (alternating 13-h temperature plateaus at 37°C and 36°C).

In vitro circadian experiments were carried out on cell populations synchronised with temperature cycles. The purpose was to determine any time dependent changes in cellular toxicity which was assessed through the measurement of cell death, cellular pharmacokinetics (drug uptake and efflux) and pharmacodynamics (drug-induced cell cycle arrest, apoptosis or autophagy (Figure II-3). Depending on the circadian experiment, e.g. cell sampling for (i) RNA extraction, aiming at the determination of circadian patterns in gene expressions or (ii) measuring cell proliferation and determining cell cycle phase distribution, measurements were taken every 4 h or 6 h in a 24 h period. In order to accommodate a 24 h circadian experiment within a 12 h working schedule, cells were subjected to antiphase temperature cycles. For example, if temperature cycle D (5-1-1-1-1) starts at 37°C, an antiphase temperature cycle would start at 32°C. The time of onset of the 12 h plateau at 37°C was defined as Time 0 (T0). Consequently, Time 12 (T12) corresponded to the start of the 12 h plateau at 32°C or 36°C. For the circadian experiments involving antiphase temperature cycles starting at 18:00, this clock hour would then correspond to T0 and T12. Sampling cells or supernatants at T0, T06, T10, T14, T18 and T22 would hence take place at clock hours 08:00 (T02 and T14), 12:00 (T06 and T18) and 16:00 (T10 and T22). Such circadian or chronopharmacology experiments were performed 2.5 days after the start of the selected temperature cycle schedule, to ensure adequate and stable circadian synchronisation of the cells in each culture dish.
Figure II-3: Common features of the experimental design of the circadian in vitro studies.

The figure depicts the timeline from the start of temperature synchronisation until the start of several in vitro studies involving drug administration as well as RNA and cell pellet sampling at different circadian time points. This was done in order to determine circadian toxicity profile of FY26, expression of clock genes, circadian inhibition of bioluminescence, time-dependent intracellular FY26 uptake and cell cycle phase distribution following FY26 administration at different circadian times.

1.6 Cytotoxicity assessment with Sulforhodamin B Assay

FY26 in vitro cytotoxicity was assessed by using the Sulforhodamin B (SRB) assay. This assay is based on the measurement of intracellular protein content (Skehan et al., 1990). Sulforhodamin B is a colorimetric assay which stains proteins by binding to protein basic amino residues depending on pH. This assay was chosen because of its advantageous sensitivity, reproducibility and linearity as compared to tetrazolium assays like MTT and XTT (Keepers et al., 1991; Rubinstein et al., 1990). It was further used to determine the IC\textsubscript{50} of FY26 both in synchronised and in non-synchronised Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 and HCT116 Per2-luc cells. The measurements were carried out using a standard protocol (Skehan et al., 1990). Four thousand Hepa1-6 cells and 8 000 HCT116 cells in 200 μL media per well were seeded irrespective of subclone in a 96-well FALCON plate (Corning, New York, USA). To
determine the IC$_{50}$ in non-synchronised cell cultures, cells were incubated for 48 h at 37°C and 5% CO$_2$ to adhere and grow. In contrast, cells on a circadian experiment underwent temperature synchronisation immediately after seeding. After 2 days for non-synchronised cells, and 2.5 days for synchronised cells, cells were exposed to the vehicle (control) or to different drug concentrations as shown in Table II-1.

*Table II-1: FY26 concentration scales used for the determination of the FY26 IC$_{50}$ concentrations.*

<table>
<thead>
<tr>
<th>Concentration Scale</th>
<th>Concentrations included (µM FY26)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>100, 10, 1, 0.1, 0.01 and 0.001</td>
</tr>
<tr>
<td>B</td>
<td>100, 10, 6, 3, 1 and 0.1</td>
</tr>
<tr>
<td>C</td>
<td>12, 10, 8, 6, 4, 2, 1 and 0.5</td>
</tr>
<tr>
<td>D</td>
<td>20, 16, 12, 8, 4, 2, 1 and 0.5</td>
</tr>
<tr>
<td>E</td>
<td>16, 12, 10, 8, 6, 4, 2, 1 and 0.5</td>
</tr>
<tr>
<td>F</td>
<td>30, 15, 10, 8, 6, 4, 2, 1 and 0.5</td>
</tr>
<tr>
<td>G</td>
<td>21, 14, 7, 6, 5, 4, 2, 1 and 0.5</td>
</tr>
<tr>
<td>H</td>
<td>3.2, 1.6, 0.8, 0.4, 0.2, 0.1, 0.05, 0.025 and 0.0125</td>
</tr>
<tr>
<td>I</td>
<td>25.6, 12.8, 6.4, 3.2, 1.6, 0.8, 0.4, 0.2 and 0.05</td>
</tr>
</tbody>
</table>

Six to 18 wells of a 96-well plate were treated with the same drug concentration or vehicle control. After completion of the 24 h drug exposure, the drug-media was removed, cells were washed with 200 µL PBS and new drug-free media was added. Cells were allowed to recover for 3 days before protein content determinations. The cells were fixed by adding 50 µL of cold 50% trichloracetic acid (TCA) (Sigma Aldrich, Gillingham, UK) directly into the media to obtain a final working concentration of 10% TCA. After 1 h of incubation at 4°C, cells were washed 10 times under slow running tap water, then air-dried. The fixed cells were treated with 50 µL of 0.4% SRB (Sigma Aldrich, Gillingham, UK) solution using 1% acetic acid (Sigma Aldrich, Gillingham, UK) as vehicle for 30 min. Cells were washed 5 times with 1% acetic acid and air dried. Two hundred µL of 10 mM Tris Base (pH 10) (Fisher Scientific, Waltham, USA) was added in each well and incubated for 1 h before
measuring the absorbance at 570 nm using a Cytation 3 (Biotek, Swindon, UK) plate reader. The obtained endpoint absorbance measurements of the colorimetric assay were normalised by setting the average of the control 100%. The data points presenting the protein content after the incubation of different FY26 concentrations were divided by the hundreds part of the average absorbance of the control. The obtained normalised data were plotted against the negative logarithms of the FY26 concentration in µM. The IC$_{50}$ was calculated as the non-linear fit of the logarithm of the concentration against the response using Graph Pad Prism 6 (GraphPad Software, Inc., La Jolla, USA).

1.7 Real-time bioluminescence recording

Bioluminescence measurement is based on the oxidation of luciferin to oxyluciferin under the contribution of luciferase, adenosine triphosphate (ATP) and oxygen (O$_2$) (Figure II-4). A by-product of this reaction is light (photons) (Buccioni et al., 2011). In order to study the synchronisation properties of dexamethasone shock, temperature cycles and the influence of drug on circadian genes expression, two different luciferase reporter with the promoter of either Per2 or Bmal1 were used (Chapter II.1.1).

![Figure II-4: Oxidation reaction of luciferin with luciferase.](image)

*This chemical reaction depicts the formation of photons as a side product of the reduction of luciferin to oxyluciferin in the presence of oxygen (O$_2$) and adenosine triphosphate (ATP) (Buccioni et al., 2011).*
To start recording real-time bioluminescence, 50,000 up to 100,000 cells harbouring a stable luciferin reporter construct were seeded in 35 mm NUNC Petri dishes containing 3 mL phenol red free and high glucose (4.5 g/mL) DMEM (Sigma Aldrich, Gillingham, UK) media. The media further contained 20 mM Hepes (Life Technologies, Carlsbad, USA), 3.75 mM L-glutamine (Prep room Life Sciences, University of Warwick), 100 μg/mL Penicillin/Streptomycin (Prep room Life Sciences, University of Warwick), 5% heat inactivated FCS and 100 μM luciferin (Promega, Madison, USA). Dishes were sealed with a thin layer of silicone grease and put into the LumiCycle (Actimetrics, Wilmette, USA). The LumiCycle is a black, closed apparatus that was put within a temperature programmable incubator which allowed to synchronise the cells (Figure II-5). The LumiCycle is equipped with four photon multiplier tubes (PMT) which are located 2.34 cm above a turntable with 32 slots, in which sealed NUNC Petri dishes can be placed. Sealing the dishes served the purpose to avoid media evaporation as humidity was set to 0% during the whole experimental duration to avoid any damage to the PMTs. The bioluminescence of each cell culture dish was measured through counting the number of photons detected by the photomultipliers over 1 min, every 10 min for a minimum of 4 days, and up to 7 days.

*Figure II-5: Experimental setup to record circadian changes in real-time bioluminescence.*
1.8 RNA extraction and Reverse Transcription

Gene expression patterns were determined by RNA extraction from cells and transcription to cDNA. The amount of gene expression was then quantified by Real-time PCR. RNA was sampled from synchronised or non-synchronised as well as from treated or non-treated cells cultured in wells of a 6-well plate (Falcon) (n = 3). Cells in a 6-well plate (Thermo Fisher Scientific, Waltham UK) were washed with 1 mL PBS and suspended in 1 mL Trizol (Thermo Fisher Scientific, Waltham, UK). The cell-Trizol suspension was transferred in a prelabelled 1.5 mL micro tube (Sarstedt, Nümbrecht, Germany). Cells were frozen at -80°C until the sampling process was finished. The RNA extraction started with defrosting the cell-Trizol samples at 20 to 22°C. Samples were kept on ice throughout the extraction and reverse transcription process to avoid degradation of the RNA. The defrosted samples were centrifuged for 1 min at 106 rcf at 4°C in order to separate cell debris from the diluted RNA, DNA and proteins. The supernatant was pipetted into a new pre-labelled tube and 200 μL of chloroform (Thermo Fisher Scientific, Waltham, UK) was added into each sample. The samples were gently mixed for 10 sec using a Stuart vortex mixer (Cole-Parmer, St. Neots, UK) and incubated for 3 min at 20 to 22°C. This step was crucial to separate DNA, RNA and proteins. The formation of the three phases was promoted by centrifugation for 15 min at 15294 rcf at 4°C. The upper transparent phase contained RNA, the middle white phase contained DNA and the lower pink phase contained the proteins. Using a 200 μL pipette (Eppendorf, Wesseling-Berzdorf, Germany), 400 μL of diluted RNA were taken out and transferred to a new prelabelled 1.5 mL micro tube. The RNA was precipitated by adding 500 μL isopropanol (Thermo Fisher Scientific, Waltham, UK) into each sample. The samples were incubated for 10 min at 20 to 22°C to ensure a complete precipitation of the RNA. The RNA was separated from the supernatant by centrifugation for 10 min at 15294 rcf and 4°C. The supernatant was carefully eliminated from the white pellet and 1 mL 75% Ethanol (VWR Chemicals, Lutterworth, UK) was added into each sample. The samples were again centrifuged for 5 min at 5974 rcf at 4°C. Ethanol was removed and the white pellet was left at 20 to 22°C to air dry. Once the white pellet has become transparent, the RNA was diluted in 15 μL diethylpyrocarbonate (DEPC, Amresco, Radnor, USA) water which was free from RNAase and DNAase. The next step involved the elimination of remaining DNA.
in the diluted RNA samples. According to the number of samples, a master mix of DNase Buffer (Promega, Madison, USA), DNase and (Promega, Madison, USA) DEPC water was prepared. Per sample, the master mix contained of 1 μL DNase Buffer, 3 μL (3U) DNase and 2 μL DEPC water. A total amount of 6 μL DNA-elimination mix together with 3 μL diluted RNA was pipetted into 0.2 mL PCR tubes (Starlab, Milton Keynes, UK) and incubated for 30 min at 37°C in order to digest the remaining DNA. The DNase was inactivated by adding 1 μL Stop Solution (Promega, Madison, USA). All samples were run on a 1% Agarose gel (Geneflow, Lichfield, UK) to confirm the success of the genomic DNA elimination. To produce the gel, 1 g Agarose was weighed in and suspended in 100 mL 1:5 TBE-DEPC water (Prep room, Life Sciences, University of Warwick). The suspension was heated up in the SHARP microwave (Scientific Laboratory Supplies, Nottingham, UK) at P80 for 2 min. Once the dilution cooled down and no bubbles sustained, the warm solution was poured into the electrophoresis plates. Whilst the gel was hardening, a mix containing 1 μL Midori Green (Geneflow, Lichfield, UK) and 8 μL DEPC water was prepared. One μL of DNA free RNA samples were added into each sample. One sample was treated with 1 μL DNA ladder (GeneDireX, Las Vegas, USA) instead of RNA. A total amount of 10 μL was pipetted into the pockets of the gel. The gel was covered with 1:5 TBE-DEPC water. The electrophoresis plates were covered and the separation of the RNA samples from possible DNA fragments was started by the exposure to 120 V and 140 A. The amount of RNA in a DNA free solution was determined with the BioDrop µLite (BioDrop, Cambridge, UK). The spectrophotometer was calibrated with 1 μL DEPC water. The amount of RNA per sample was then determined by using 1 μL per sample which was pipetted on the sensor. The amount of RNA was measured in ng/μL. Additionally, the ratio of A260/A230 and A260/280 representing the phenol/chloroform and protein/background purity was determined respectively. Based on the RNA amount per sample, DEPC water was added to generate a final concentration of 200 ng/μL in each sample. In order to start the Reverse Transcription, a master mix of 1 μL (200 ng) Random Primers (Thermo Fisher Scientific, Waltham, UK), 1 μL 10 mM dNTP Mix consisting of equal volumes of 100 mM dATP, dCTP, dGTP and dTTP (VWR, UK) and 8 μL DEPC water per sample was prepared according to the total number of samples. Per sample, 10 μL of the master mix and 2 μL of RNA (400 ng/μL total RNA
concentration) were pipetted into the new 0.2 mL PCR tubes. The mixture was heated up for 5 min at 65°C. Per sample, 6 μL containing 2 μL 0.1 M DTT (Invitrogen, Waltham, USA) and 4 μL 5X First-Standard Buffer (Invitrogen, Waltham, USA) were added. The content per tube was gently mixed and incubated for 2 min at 25°C. The reverse transcription reaction was initiated by transferring 1 μL (200 U) of Superscript II RT (Invitrogen, Waltham, USA) into each tube. The content was homogenised and then incubated in the Q-Cycler 96 (HAIN Technology, Byfleet, UK) for 10 min at 25°C, 50 min at 42°C and 15 min at 70°C to stimulate reverse transcription. Once the temperature cycle was finished, the cDNA was stored until further use at -20°C.

1.9 Real-time Polymerase Chain Reaction (PCR)

The previously obtained cDNA was used to determine circadian clock gene expression. Therefore, Real-time PCR (RT-PCR or qPCR for quantitative PCR) was applied to determine the expression of the clock genes *Bmal1*, *Per2* and *Rev-erba*. 36B4 was chosen as a reference housekeeping gene as its expression remained stable (Ramanathan *et al.*, 2012; Zhang *et al.*, 2016). Three technical replicates were taken from each of the 3 biological cDNA replicates. Thus, 9 samples corresponded to each experimental condition. Based on the number of samples, a reaction mix was prepared. The reaction mix for one sample was composed of 0.4 μL forward primer 0.4 μL reverse primer (Integrated DNA Technologies BVBA, Leuven, Belgium), and 3.2 μL purified water from PURELAB® flex (ELGA Veolia, High Wycombe, UK) and 5 μL 2X SensiFAST SYBER No-ROX Mix (Bioline, London, UK). Each gene required a different reaction mix since one reaction mix contained primer for only one gene. Thus, 5 different reaction mixes were prepared containing the forward and reverse primer of the genes *Bmal1*, *Per2*, *Rev-erba*, *LC3B* and 36B4 (Table II-2). In a 96-well plate or 384-well plate (AXYGEN Scientific, New York, USA), 9 μL of reaction mix was pipetted in each well followed by the addition of 1 μL cDNA. The plate was sealed using MicroAmp® Clear Adhesive Film (Life Technologies, Carlsbad, USA) and put in the FluoroCycler® 96 (HAIN Technologies, Byfleet, UK). The samples were subjected to the preprogrammed temperature cycles consisting of an initial heating up phase for 2 min to 95°C. This was followed by a cycle at 95°C for 6 min and 60°C for 25 min. This second cycle was repeated in total 40 times before the melting curve was initiated to test whether single and specific DNA products have been produced.
Table II-2: Forward (F) and reverse (R) sequences of mouse (m) clock gene and autophagy primers.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>m36B4-F</td>
<td>5′ GCT GAT GGG CAA GAA CAC CA′-3</td>
</tr>
<tr>
<td>m36B4-R</td>
<td>5′ CCC AAA GCCTGG AAG AAG GA′-3</td>
</tr>
<tr>
<td>mBmal1-F</td>
<td>5′ GGC TGG ACG AAG ACA ATG AGC′-3</td>
</tr>
<tr>
<td>mBmal1-R</td>
<td>5′ GTT GTG GAA CCA TGT GCG AGT′-3</td>
</tr>
<tr>
<td>mPer2-F</td>
<td>5′ AGA GCA CAG GGT CTG GAG GA′-3</td>
</tr>
<tr>
<td>mPer2-R</td>
<td>5′ TGG AAC ACA GGT AGG GGG TAA′-3</td>
</tr>
<tr>
<td>mRev-erbα-F</td>
<td>5′ GGC TGA TTC TTC ACA CAC ACA C′-3</td>
</tr>
<tr>
<td>mRev-erbα-R</td>
<td>5′ GGT CTT GGG GTG GCT ATA CTG′-3</td>
</tr>
<tr>
<td>mLCl3B-F</td>
<td>5′ CCC ACC AAG ATC CCA GTG AT′-3</td>
</tr>
<tr>
<td>mLCl3B-R</td>
<td>5′ CCA GGA ACT TGG TCT TGT CCA′-3</td>
</tr>
</tbody>
</table>

1.10 Time dependent pharmacodynamics endpoints of FY26

1.10.1 Effect of FY26 on apoptosis

In order to test the effect of FY26 on the induction of apoptosis, Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 and HCT116 Per2-luc cells from each well of a 6-well FALCON plate (Corning, Liversedge, UK) were incubated with different FY26 concentrations (5 µM, 7.5 µM and 10 µM FY26) 2.5 days after the start of temperature synchronisation. The cell pellet sampling took place 24 h after the start of drug exposure. Cell pellets were sampled every 4 h in a 12 h time span using antiphase temperature cycles of the temperature program D as described in the previous chapter II.1.5.3. The whole media content in each well was transferred into a 15 mL tube to collect the dead cells (Sarstedt, Nümbrecht, Germany). Adherent cells in each well were washed with 1 mL PBS. PBS as well as media were pipetted into the 15 mL tube. Cells were treated with 300 µL Trypsin-EDTA. Trypsinisation was stopped by adding 700 µL fresh media followed by the collection of the whole 1 mL cell suspension into the 15 mL tube. Cells were centrifuged at 200 rcf for 5 min at 22°C to separate the cells from the media. The supernatant was removed and the cell pellet was suspended with 5 mL PBS. In order to add the staining solution, PBS was removed from the cells.
Methods and Materials

after centrifugation (200 rcf, 5 min at 22°C). Each cell pellet was suspended in 100 μL staining solution composed of 2.5 μL of a 1 mg/mL Propidium iodide solution (PI, Cambridge Bioscience LTD, Cambridge, UK), 2.5 μL Annexin-V (Biolegend, San Diego, USA) and 95 μL 1X Binding Buffer (Biolegend, San Diego, USA), and transferred into a FALCON polystyrene flow cytometry tube (Corning, Liversedge, UK). Three controls were required, of which one was stained only with Annexin-V (5 μL Annexin, 195 μL 1X Binding Buffer) and a second one with PI (5 μL PI and 195 μL 1X Binding Buffer). The third control was left unstained and diluted in 200 μL 1X Binding Buffer only. After a 30 min incubation time in the dark at 22°C, the respective total numbers of apoptotic, necrotic and viable cells were determined by exposing cells to different lasers using a flow cytometer (Fortessa, BD Bioscience, Berkshire, UK). A yellow laser (YG561-610/20 nm) and a blue laser (B488-530 nm) were chosen to detect PI and Annexin-V respectively at 200 V. PI is able to penetrate the cell membrane of dead cells due to the loss of membrane integrity. Annexin-V is characterised by a high affinity towards phosphatidylserine. As the cell undergoes apoptosis, phosphatidylserine is translocated from the inner to the outer cell membrane resulting in Annexin-V binding. Through use of PI and Annexin-V in combination, apoptotic (Annexin-V positive, PI negative), necrotic (Annexin-V positive, PI positive) and viable cells (Annexin-V and PI negative) were detected due to a loss of membrane integrity according the cellular stage (Koopman et al., 1994; Vermes et al., 1995). The Flowing Software 2.5.1 (by Perttu Terho, University of Turku, Finland) was used for data analysis.

1.10.2 Time dependent effect of FY26 on cell cycle phase distribution

Sample collection and preparation matched the description in 2.12 until the staining step. The cell pellet was suspended in 1 mL ice cold 70% ethanol and incubated for at least 1 h at 2°C to 8°C in order to fix them. Cells were allowed to stay fixed in 70% ethanol for up to 24 h until the sample collection at different time points was finished. Once all the cells were collected, the sample preparation took place simultaneously. Therefore, ethanol was separated from the cells through centrifugation at 200 rcf for 5 min at 22°C and removed. The cell pellet was transferred into a polystyrene tube with 500 μL staining mix composed of 25 μL 1 mg/mL PI, 50 μL 100 mg/mL RNase (VWR, Lutherworth, UK) and 425 μL PBS. After 30 min of incubation, the cells were
separated from the staining solution by centrifugation at 200 rcf for 5 min and suspended in 500 μL PBS. One replicate of the control was left unstained to better adjust the laser settings for the measurement with the flow cytometer (Fortessa flow cytometer, BD Biosciences, Wokingham, UK). A yellow laser (YG651-610/20 nm) with a voltage of 200 V was used during the whole measurements. The gating of the different cell cycle phases, G0/G1-phase, S-phase and G2/M-phase, was done with the free flow cytometry data analysis software “Flowing Software 2.5.1” developed by Perttu Terho (University of Turku, Finland). The gates were set using controls which were stained or not with PI in order to set a template for the gates. This template was subjected to all treated controls. The cell number was acquired by the software and further expressed as relative number of cells per phase normalised to the non-treated controls.

1.11 Circadian Rhythm-Dependent in vitro Pharmacokinetics of FY26

1.11.1 Aim of the study

This study aimed to determine whether cellular chronopharmacokinetics would drive circadian patterns in in vitro toxicity. The main focus was the measurement of the time dependent cellular uptake, intracellular concentrations, and elimination of FY26 over the 72 h following initial drug exposure. The study further assessed circadian changes in the time (t_{max}) and quantity (C_{max}) of maximum drug uptake, and subsequent intracellular and extracellular pharmacokinetics.

1.11.2 Experimental design

Fifty thousand Hepa1-6 Per2-luc cells per 3 mL media were seeded in each of 3 wells of a 6-well FALCON plate. In total, 60 6-well FALCON plates were prepared. Each of them corresponded to a different experimental condition. After seeding, half of the plates were put in a first temperature programmable incubator and the other half in the second incubator. The temperature cycle program D in the first incubator was in antiphase to the second one thus starting at 37°C in incubator 1 and at 32°C in incubator 2 (Figure II-6). After 2.5 days, the media was replaced by media containing 1 μM FY26 for all the dishes except for the six “baseline” samples labelled as “0 h”
which served as FY26-free controls. Cells were treated with 1 µM FY26 starting 2 h, 6 h and 10 h after the last temperature switches (i) from 36 to 37°C after 2.5 days, corresponding to T02, T06 and T10 and (ii) from 37 to 36°C, corresponding to T14, T18 and T22. Following FY26 exposure at each circadian time point (T02, T06, T10, T14, T18 and T22), cells and media from 3 wells (triplicates of each experimental condition) of a 6-well FALCON plate were harvested 1 h, 3 h, 9 h, 12 h, 18 h, 24 h, 30 h, 36 h, 48 h and 72 h after drug exposure in order to determine the in vitro chronopharmacokinetics of FY26 through the measurement of extracellular (media) and intracellular (cells) osmium concentrations (Figure II-6 and Figure II-7). Importantly, cells were exposed to 1 µM FY26 throughout the experiment.

**Figure II-6: Experimental design of time dependent intracellular FY26 uptake.**

*Hepa1-6Per2-luc cells were synchronised to antiphase 24 h temperature cycles with an initial 5°C difference followed by cycles with a 1°C difference with a start at 37°C and a 12 h later decrease to 32°C for incubator 1 and start at 32°C with an increase to 37°C for incubator 2. Cells were kept under synchronisation for the total experimental duration. After 2.5 days of synchronisation, cells were exposed to 1 µM FY26 at T02, T06, T10, T14, T18 and T22 with 3 replicates per condition.*

With regard to the extracellular osmium concentration, 1 mL from each of 3 wells in a 6-well FALCON plate was transferred into a labelled 1.5 mL micro tube and centrifuged at 10 621 rcf for 4 min. With a 1 000 µL pipette (Eppendorf, Wesseling-Berzdorf, Germany), 900 µL of the media supernatant were taken out and relocated into a new 1.5 mL micro tube. Those samples were stored at -80°C. The remaining media in each well was removed and cells were washed with 2 mL PBS in order to
determine the intracellular osmium content. After the removal of PBS, cells were incubated with 300 µL Trypsin-EDTA for 3 min. The trypsinisation process was stopped by adding 700 µL fresh media. The cells were counted. The 1 mL media cell suspension was pipette into a 1.5 mL micro tube and cells were separated from the media through centrifugation at 10 621 rcf for 4 min. The supernatant was removed and the cell pellet was frozen at -80°C as well as the media sample for the measurement of osmium content with inductively-coupled plasma-mass spectrometry (ICP-MS).

In order to ensure the collection of each cell pellet and media sample, I was kindly offered help from Doctor Swati Kumar. She sampled the cell pellets and media samples on Day 2 from 11:00 to 23:00.

![Figure II-7](image)

**Figure II-7:** Time schedule of media and cell pellet sampling referred to circadian time of drug addition, with corresponding clock hours and days of the experiment.

After drug addition at T02, T06, T10, T14, T18 and T22 cells were harvested and media samples were taken at 1 h, 3 h, 9 h, 12 h, 18 h, 24 h, 30 h, 48 h and 72 h after the beginning of drug exposure. Cell pellets and media samples were stored at -80°C for ICP-MS.

### 1.11.3 Drug content determination through inductively-coupled plasma-mass spectrometry

In a next step, the osmium concentration in digested cells (cell pellet) and in media was determined. Cellular osmium content (189Os) was measured by incubating each cell pellet with freshly distilled 73% nitric acid and double deionized water overnight at 80°C. After the incubation time was finished, the resulting solutions were diluted with thiourea and ascorbic acid to a working concentration of 3.6% v/v nitric acid,
10 mM thiourea and 0.1 g/L ascorbic acid. The addition of thiourea and ascorbic acid served the purpose to stabilise osmium and to prevent oxidation to OsO$_4$(Venzago et al., 2013). The sample preparation to determine the osmium concentration in the cell media was done by diluting 50 µL cell media with 4950 µL of stock solution containing 10 mM thiourea, 0.1 g/L ascorbic acid in 3.5% v/v nitric acid. The concentration of osmium in the cell and in the cell media was determined by ICP-MS (Agilent Technologies 7500 Series ICP-MS, Cheshire, UK) in no gas mode. Before the sample measurement, the ICP-MS was calibrated using calibration standards ranging from 0.1 to 100 ppm diluted in a solution of 36% v/v nitric acid, 10 mM thiourea and 0.1 g/L ascorbic acid. The isotope of erbium ($^{167}$Er) was used as internal standard. Data acquisition and analysis were carried out using ICP-MS B.03.06 software (Agilent Technologies, Cheshire, UK). The amount of osmium was obtained in parts per million (ppm) and normalised to nanogram per 1 million cells (ng Os/ million cells).

Doctor Carlos Cano-Sanchez (Department of Chemistry, University of Warwick, and Coventry, UK) and myself prepared the samples and determined the osmium concentration with the ICP-MS.
2 In Vivo Methodology

2.1 Animals and housing

In vivo studies were carried out using adult “wildtype” or PER2::LUC transgenic C57BL/6 mice, or CD1-\textit{Foxn1}\textsuperscript{nu} mice. Male C57BL/6 mice aged 5 to 9 weeks were used for the determination of circadian toxicity patterns of FY26 or FY25. The PER2::LUC C57BL/6 mouse served the determination of FY26 toxicity and its effect on the PER2 expression and core body temperature. This mouse was initially engineered by the laboratory team of Joseph Takahashi (UT Southwestern Medical Center, Dallas, SA) (Yoo et al., 2004). The luciferase gene was knocked in the sequence of clock gene \textit{Per2} after the terminal exon of the endogenous m\textit{Per2} locus resulting in the expression and translation of a PER2::LUCIFERASE fusion protein. It has been shown that PER2::LUC fusion protein is expressed in the SCN, pituitary, lung, tail, kidney and liver with highest expressions in the SCN and in liver. No phenotypic difference was shown between C57BL/6 wildtype mice and C57BL/6 PER2::LUC mice (Yoo et al., 2004). The CD1-\textit{Foxn1}\textsuperscript{nu} mice are athymic immunodeficient nude mice on CD1 background. These male mice were implanted with 1 \cdot 10^6 Hepa1-6 \textit{Per2-luc} cells in a 100 \mu L suspensions at 9 to 15 weeks of age, in order to investigate FY26 efficacy at different dosing times, in the absence of immunologic rejection of the cancer cells.

C57BL/6 mice were purchased from Charles River (London). PER2::LUC mice were kindly gifted by Doctor Xiao Mei Li, from INSERM U935 (Villejuif, France) and bred in our animal facility. The CD1-\textit{Foxn1}\textsuperscript{nu} mice were either purchased from Charles River (London) or bred in the animal facility at Warwick University. A single up to 4 mice were kept in Techniplast Blue Line 1284L individually ventilated cages (IVC) (Techniplast, Leicester, UK) under positive pressure and permanent ventilation in the animal facility unless otherwise specified. Mice had access to food and water ad libitum.
2.2 Entrainment of mice using Light-Dark Cycles with light onsets at different real-times

The purpose of the entrainment of mice to different light-dark cycles was to fit a 24 h circadian experiment into a 12 h working day. This was achieved by exposing mice to regular alternations of light (L) for 12 h and darkness (D) for 12 h (LD12:12) with light onset and offset occurring at different clock hours. Light onset was defined as Zeitgeber Time 0 (ZT0), and dark onset as ZT12 under LD12:12 entrainment. For example, in order to treat a first group of mice at ZT06 and a second one at ZT18 at the same clock hour 13:00, light onsets were programmed at clock hours 07:00 and 19:00 respectively (Figure II-8 B). The entrainment was achieved using a dedicated chronobiological cabinets (Eurobioconcept, Vitry, France) consisting of 5 compartments each (Figure II-8 A). For each compartment a different light-dark cycle can be programmed. Each chronobiological cabinet is equipped with a system that provides adjustable continuous air flow through filters to each compartment, which are thus exposed to minimal environmental contamination and fresh air. Temperature and light intensity were automatically recorded through sensors in each compartment that were transmitting data wireless to a dedicated computer every 10 sec throughout the whole experimental duration starting with the beginning of the light-dark entrainment. Environmental temperature in the animal facility was kept between 20 and 24°C. The light intensity in the cabinets varied between 100 and 160 lux at cage level. The light source was a 300*1200 LED Panel CW/DW/WW (Eurobioconcept, Vitry, France) with 4200Lm/5200Lm. Room humidity was monitored daily by a humidity meter installed and kept between 45 and 65% respectively. All the mice had free access to food and water throughout each experiment.
Figure II-8: Experimental set up of mice entrainment to LD 12:12 cycles.

(A) Chronobiologic cabinet consisting of 5 compartments allowing the programming of different light dark cycles with a different light on and off set. (B) Example for an antiphase entrainment schedule using two compartments.

2.3 Drug preparation

The anticancer drug candidates FY26 and FY25 were synthesised and purified (> 98% purity) in the laboratory of Professor Peter J. Sadler in the Chemistry Department of the University of Warwick (Fu et al., 2010). Throughout the different studies, mice were injected with 20 mg/kg, 30 mg/kg, 40 mg/kg, 50 mg/kg, 60 mg/kg, 80 mg/kg or 160 mg/kg of FY26 and/or FY25. Both drugs were diluted in the vehicle, composed of 1% TWEEN 80 (Sigma Aldrich, Gillingham, UK), 5% DMSO (Corning, New York, USA) and 94% of 0.9% Saline (Prep room, Life Sciences, University of Warwick). The drug preparation was done under laminar flow guaranteeing aseptic conditions. Each mouse was injected with a volume of 10 µL drug-vehicle dilution per gram body weight. The dose was confirmed following drug preparation by Doctor Carlos Cano-Sanchez using ICP-MS as described previously.

With regard to the drug preparation, it was essential to know the number of mice receiving treatment as well as the average mouse body weight. Based on these data, the quantity of drug needed was calculated. The calculation for the treatment of 35
Mice with 50 mg/kg FY26 are explained in Figure II-9. This method required to determine the body weight of each mouse before injection in order to ensure the injection of the precise volume and dose. All injections were administered intraperitoneally (i.p.) using 1 mL sterile Plastipak™ syringes and sterile 0.5 and 16 mm Microlance™ 3 needles (BD, Wokingham, UK).

### 1. Amount of drug
- $50\,\text{mg kg}^{-1} = 0.05\,\text{mg g}^{-1}$
- $0.05\,\text{mg g}^{-1} \times 30\,\text{g} = 1.5\,\text{mg}$

A mouse with 30 g body weight received 1.5 mg FY26 which equals a dose of 50 mg/kg.

- $1.5\,\text{mg} \times 36 = 54\,\text{mg}$

54 mg FY26 is needed to treat 36 mice with a single dose of 50 mg/kg.

### 2. Volume of vehicle
- $30\,\text{g} = 300\,\mu\text{l}$

A mouse with 30 g body weight will be injected with a total volume of 300 µl.

- $1.5\,\text{mg} = 300\,\mu\text{l} \rightarrow 0.05\,\text{mg} = 10\,\mu\text{l}$
- $\frac{54\,\text{mg}}{0.05\,\text{mg}} = 1080 \rightarrow 1080 \times 10\,\mu\text{l} = 10.8\,\text{ml}$

54 mg FY26 were diluted in 10.8 ml vehicle to ensure a treatment with 50 mg/kg.

*Figure II-9: Example of FY26 dose calculation for in vivo studies.*

### 2.4 Humane endpoints

Throughout the *in vivo* experiments, any mouse showing signs of pain or distress exceeding the severity limit ‘moderate’ was culled. Body weight loss was used as a marker for mouse welfare. A mouse with a body weight loss of 20% or more within 24 h or a persistent body weight loss of 15% over 72 h without recovery was euthanized. Additionally, tumour-bearing mice were allowed to have a maximum tumour weight of 2 g. Mice with a tumour weight exceeding those limits, were sacrificed during the study but data obtained from these mice were still included in the study. All these regulations applied at any time and were followed regardless of the success of the ongoing experiments.
2.5 Overview of in vivo studies

Five mouse studies aimed to respectively determine the maximum tolerated dose (MTD) of FY26, the time dependent changes in the toxicity of this FY26 and FY25, the effects of FY26 on biomarkers of the CTS, including circadian rhythms in core body temperature and liver PER2 expression. The fifth study assessed the antitumour efficacy of FY26 and preliminary time dependent changes in tumour growths inhibition (Table II-3). The specific design details are explained in the following section.
Table II-3: Overview of the in vivo studies (N=number)

<table>
<thead>
<tr>
<th>Study goal</th>
<th>Specific objective</th>
<th>Mouse strain (sex)</th>
<th>Main endpoints</th>
<th>Total N of mice</th>
<th>N of mice per experiment</th>
<th>N of experiment</th>
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<tbody>
<tr>
<td>Dose Finding</td>
<td>Determination of the MDT of FY26 and FY25</td>
<td>C57BL/6 mice (male)</td>
<td>Body weight change</td>
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<td>6</td>
<td>2</td>
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<tr>
<td>Chronotoxicity</td>
<td>Determination of circadian toxicity profile of FY26 and FY25, and identification of the circadian time points of best and worst tolerability</td>
<td>C57BL/6 mice (male)</td>
<td>Body weight change, Survival</td>
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<td>78</td>
<td>3</td>
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<tr>
<td>Histopathology</td>
<td>Determination of FY26-induced tissue lesions</td>
<td>PER2::LUC C57BL/6 mice (male)</td>
<td>Histologic lesions, apoptosis, necrosis tissue alterations</td>
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<td>6</td>
<td>1</td>
</tr>
<tr>
<td>CTS effects on temperature</td>
<td>Determination of FY26 effects on core body temperature rhythm following dosing at ZT06 or ZT18.</td>
<td>C57BL/6 mice (male)</td>
<td>Core body temperature rhythm and body weight changes</td>
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<tr>
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<td>Description</td>
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<td><strong>CTS effects on the liver clock</strong></td>
<td>Simultaneous determination of circadian patterns in PER2 liver expression and core body temperature following dosing at ZT06 or ZT18. PER2::LUC C57BL/6 mice (male)</td>
<td>Circadian rhythms in PER2 expression, and core body temperature, and changes in body weight</td>
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<td>3</td>
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<td><strong>Tumour growth inhibition</strong></td>
<td>Determination of the therapeutic FY26 dose needed to inhibit tumour growth. CD1-Foxn1nu mice (male)</td>
<td>Tumour and body weight</td>
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<tr>
<td><strong>Time dependent tumour growth inhibition</strong></td>
<td>Preliminary determination of tumour growth inhibition following FY26 dosing at ZT06 or ZT18. CD1-Foxn1nu mice (male)</td>
<td>Tumour and body weight</td>
<td>20</td>
<td>20</td>
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2.6 Dose finding study

2.6.1 Aim
This study was conducted in order to determine the maximum tolerated dose level of FY26 and FY25, to be selected to further identify any circadian toxicity pattern in a follow up experiment. The requirement for the optimal dose levels was a body weight loss between 15 to 20%.

2.6.2 Experimental design
Twelve male C57BL/6 mice aged 5 to 6 weeks were housed in 4 cages containing 3 mice each. Mice were used for 2 consecutive experiments and kept in the common animal facility with lights on at 07:00 and lights off at 19:00. Each dose was given as a single intraperitoneal (i.p.) injection. Eight to 9 weeks old mice in cages 1 and 2 were injected with vehicle (control) or FY26 (30 mg/kg, three mice). The body weights of all mice were measured 6 h after injection and daily for 5 days thereafter. On day 5, the same mice received a second i.p. injection of vehicle or FY26 (40 mg/kg). Body weight was measured as previously mentioned daily for 7 days. A further experiment involved the i.p. administration 40 mg/kg of FY26 to the 3 mice in cage 3 and 60 mg/kg of FY26 to the 3 mice in cage 4, now aged 10-11 weeks, and subsequent body weight determinations as per previous protocol for 5 days.

2.7 Circadian toxicity of FY26 and FY25

2.7.1 Aim
The purpose of this study was to investigate whether FY25 and FY26 toxicities would display a circadian pattern in vivo. The study involved three consecutive experiments with a similar design. Experiment (Exp) II and III had identical designs, whereas the design of Exp I differed slightly. First the common features and secondly the specificities of the Exp are outlined.

2.7.2 Common features of experimental designs
Two hundred-thirty four male C57BL/6 mice aged 5 to 9 weeks were purchased from Charles River (London, UK). Mice were allocated into cages hosting 1 to 3 mice per
cage. They were allowed to adjust for one week to the common animal facility before the start of light-dark entrainment in the dedicated chronobiological cabinets. Mice were distributed amongst 18 groups. Six groups of 18 mice each per drug and time point were injected with 50 mg/kg of FY25 or FY26 at ZT02, ZT06, ZT10, ZT14, ZT18 or ZT22. Three mice in each of six control groups were injected with vehicle only at the above-mentioned circadian time points. All the mice were aged 11 to 12 weeks at the time of treatment administration. Mice were weighed twice a week before injection, and daily thereafter. Moreover, they were visually inspected every day following treatment.

2.7.3 Specific aspects of each experiment design

Specific aspects were related to the age of the mice upon purchasing, the time left for adjustment to the common animal facility, the allocation of mice to cages and groups, and the duration and schedules of light-dark entrainment.

Mice arrived from the breeder at ages 5 to 6 weeks for Exp I, and 8-9 weeks for Exp II & II. Mice in the first Exp were allowed to adjust to the common animal facility for two weeks followed by an entrainment period of 30 days. The adjustment time of mice from Exp II & III was only 5 days long in the common animal facility before the start of the light-dark entrainment for 21 days.

A. Treatment plan Experiment I

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B. Treatment plan Experiment II & III

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Figure II-10: Experimental differences in the allocation of mice in groups.
This figure presents the distribution of mice per treatment group and emphasises that in Exp I the control mice were entrained individually in two additional compartments whereas in Exp II & III all mice injected at one time point were entrained to the same light dark cycle.

Regarding the allocation of mice, 8 compartments of the chronobiological cabinets were used. The light dark cycles in the first 6 compartments were programmed with a lights on at clock hours 05:30 to 17:30 (for ZT02 treatment, corresponding to 07:30), 03:30 to 15:30 (ZT06), 01:00 to 13:00 (ZT10), 23:30 to 11:30 (ZT14), 21:00 to 09:00 (ZT18), 19:00 to 07:00 (ZT22), 08:00 to 20:00 (Control ZT02, ZT06, ZT10) and 04:30 to 16:30 (control ZT14, ZT18, ZT22). Each of the compartments contained 4 cages with 3 mice of which half of the mice were treated with 50 mg/kg FY25 and the other half with 50 mg/kg FY26. The circadian treatment times ZT02, ZT06, ZT10, ZT14, ZT18 and ZT22 corresponded to clock hours 07:30, 09:30, 11:00, 13:30, 15:00 and 17:00 respectively. The last 2 compartments contained three cages with one mouse per cage serving as the vehicle control. Those mice were injected with the vehicle at clock hours 10:00 (ZT02), 10:30 (ZT14), 14:00 (ZT06), 14:30 (ZT19), 17:00 (ZT22) and 18:00 (ZT10) (Figure II-10 and Figure II-11).
**Figure II-11: Entrainment protocol of Exp I.**

Eight light dark cycles were programmed enabling a treatment of 6 mice each with 50 mg/kg FY26 and FY25 at the circadian time points ZT02, ZT06, ZT10, ZT14, ZT18 and ZT22 in a 12 h time span. The 6 control mice were programmed to 2 additional light dark cycles different from the mice treated with FY26 and FY25.

Compared to Exp II & III, only 6 compartments of the chronobiological cabinet were used of which each compartment was programmed with a different light dark cycles with a light on and off set at 08:00 to 20:00 (ZT02), 04:30 to 16:30 (ZT06), 01:00 to 13:00 (ZT10), 23:00 to 11:00 (ZT14), 19:30 to 07:30 (ZT18) and 16:00 to 04:00 (ZT22). Treatment was thus possible at clock hour 10:00 (ZT02), 10:30 (ZT06), 11:00 (ZT10), 13:00 (ZT14), 13:30 (ZT18) and 14:00 (ZT22). Each compartment hosted 13 mice allocated in 5 cages of which 3 cages contained 3 mice and 2 cages 2 mice. Six
of the mice received treatment with 50 mg/kg FY25, another 6 mice with 50 mg/kg FY26 and the 13\textsuperscript{th} mouse was injected with the vehicle (Figure II-10, Figure II-12).

![Entrainment schedule Exp II & III](image)

**Figure II-12: Entrainment schedule of Exp II & III to access circadian toxicity.**

Six light-dark cycles were programmed to enable treatment of each mouse at ZT02, ZT06, ZT10, ZT14, ZT18 and ZT22 during daytime hours.

### 2.8 Histopathology Study

#### 2.8.1 Experimental design

Six male PER2::LUC C57Bl/6 mice, aged 12 to 15 weeks, were kept in LD12:12 with L from 07:00 to 19:00. Mice were treated with 4 intraperitoneal injections of FY26 (50 mg/kg i.p., 4 mice) or vehicle (2 mice) at ZT06 every other day. Mice were weighed and visually inspected daily. Twenty-four hours after the fourth injection, mice were sacrificed, and liver, both kidneys, stomach, ileum, colon and the sternum were excised, washed in PBS and fixed in 10% Neutral Buffered Formalin (CellPath
LTD, Powys, UK). After 24 h in formalin, organs were transferred into PBS and send for slicing to the Histology Facility, University of Manchester (Mr Peter Walker). Histopathological lesions were kindly assessed by Dr Kishore Gopalakrishnan (University Hospital Coventry and Warwickshire, UK).

2.9 Effect of FY26 administration on core body temperature

2.9.1 Aim

This was designed in order to assess the effect of FY26 on mice core body temperature representing one of the outputs of the circadian timing system.

2.9.2 Implantation of the Anipill® and intraperitoneal temperature recording

The continuous recording of intraperitoneal temperature involved the use of an implanted electronic capsule, equipped with a temperature sensor and tele-emitter (Anipill®, Bodycap, Paris, France). The Anipill®, a 1.77 cm long and 0.89 cm wide capsule with a weight of 1.7 g, measured core body temperature every 5 or 15 min according to the investigator’s selection. The stored measurements were transmitted to the data receiver, the Aniview Monitor® (Bodycap, Paris, France). The Aniview Monitor® was then connected to a computer for data uploads, and their organisation and visualisation in the Anipill Software V2 (Bodycap, Paris, France). The data files were then transferred into Excel formats for further analyses. In this experiment, core body temperature was measured every 5 min, and data were tele-transmitted through placing the monitor within 1 meter of the Anipill®-implanted mouse. The data synchronisation between the Anipill® and the Aniview Monitor® took place every other day (Figure II-13).

The capsule was surgically implanted into six male C57BL/6 mice aged 9 to 10 weeks, before the start of circadian entrainment. Surgery tools were sterilised. Each Anipill® was activated using the Aniview Monitor® and the Anipill® activator device by placing the Anipill in the activator with the monitor next to it. The activation was initiated by pushing the “Start” button. Afterwards, the Anipill® was washed with mild detergent and sterilised with disinfectant (CIDEX PLUS®) for at least 10 h. On the surgery day, the Anipill® was taken out from the disinfectant solution and placed in a 10 cm Petri dish (Sarstedt, Germany) filled with sterile saline solution. Prior to implantation, each
mouse was injected with 100 µL of a solution of 0.01 mg/mL of buprenorphine (0.3 mg/mL Vetergesic, Bonaventura Locums LTD, UK) and 100 µL of a solution of 0.04 mg/mL of meloxicam (0.5 mg/mL Metacam, Bonaventura Locums LTD, UK). The mouse was introduced to 1.5 to 3.5% isoflurane with an oxygen flow rate of 2 L/min. The skin area to be operated was cleaned and a straight longitudinal incision of 1 to 2 cm of the skin, then the peritoneum was made using a sterile razor blade. The skin was separated from the abdominal muscle wall using the round tip of scissors. To open the abdomen, the muscle was held up with tweezers and a small horizontal 1 to 2 cm cut was made. With sterile tweezer, the Anipill® was positioned into the left side of the peritoneal cavity. The peritoneum was then sutured, the skin moisturised with 0.9% saline and closed with 3 to 4 wound clips. The mouse was allowed to recover in a cage situated on a heated mat.

![Figure II-13: Experimental procedure to measure core body temperature in mice.](anipill_monitor.png)

Mice were inspected 1, 2 and 4 h after surgery, then visually checked and weighed daily for 7 days. The staples were removed 7 to 14 days after surgery.

### 2.9.3 Experimental design after implantation

The mice were randomly allocated to 2 groups of 3 mice each and subjected to LD 12:12, with L from 08:00 to 20:00 for Group 1, and from 20:00 to 08:00 for Group 2 (Figure II-14). Mice were single housed with access to water and food ad libitum. Mice were kept under entrainment, while core body temperature was measured every 5 min and automatically stored in the capsule’s memory throughout the experimental duration. They were injected intraperitoneally with the vehicle on day 34, then with FY26 (50 mg/kg i.p.) on day 43, and euthanized on day 55. The data transmission...
followed a 5 min synchronisation process between capsule and data receiver. The Anipill® was removed post mortem from the abdomen of each mouse, cleaned and deactivated.

![Experimental set up of the determination of the effect of FY26 on the circadian clock using body temperature rhythm as a circadian biomarker.](image)

**2.10 Effect of FY26 on PER2::LUC liver expression and core body temperature**

**2.10.1 Aim**

This study aimed to determine the effect of FY26 on the circadian timing system and on the molecular clock, using the PER2::LUC mouse model. Towards this goal, liver PER2::LUC expression and core body temperature 24-h patterns were jointly measured at baseline and after a single i.p. injection of 50 mg/kg of FY26.

**2.10.2 Experimental design**

Nine PER2::LUC mice, aged 9 to 11 weeks, were entrained to antiphase LD12:12 cycles, with L (light) onset at 07:00 for four mice, or at 19:00 for the other five animals. Treatment was scheduled at ZT06 and ZT18, corresponding to clock hour 13:00. The study consisted of 3 experiments, each of them including one mouse in a given LD
12:12 schedule, and two mice in an antiphase LD 12:12 regime or the reverse. The start of entrainment for Exp I was scheduled 2 weeks before the start of Exp II and 4 weeks before the start of Exp III. Each group experienced a total entrainment period of 3 weeks which was split into 2 weeks of entrainment in the chronobiologic cabinets and one week of entrainment in the Real-time Biolumicorder (RT-BIO, LESA Technology, and Geneva, Switzerland) with the implantation of the Anipill® occurring 2 weeks after the start of the LD 12:12 entrainment. Three RT-BIOS were kindly loaned by INSERM-Villejuif (France) to the Chronotherapy Team at WMS. The RT-BIO is equipped with a bioluminescence measuring system engineered for the continuous, non-invasive and high frequency measurements of bioluminescent gene expression in freely moving mice for weeks, thus enabling in vivo molecular circadian dynamics investigations. It consists of a mouse cylindrical metal cage, with a photon multiplier situated on the top of the cage, as well as a food hopper and water reservoir to ensure food and water supply ad libitum (Saini et al., 2013). The cylindrical cage is closed by a second cylindrical lead to ensure a complete isolation from external light sources. Through a ventilation system, air exchange was guaranteed throughout the experimental duration (Figure II-15). In order to study molecular clock function in vivo, transgenic mice expressing the fusion protein PER2::LUC were used. A water solution containing 0.78 mg/mL of luciferin filled the cage bottles. Since fur prevents the emission of photons, the area of the right lower back of the mice was shaved so as to predominantly count the photons emitted by the liver (Saini et al., 2013). This method allowed to investigate the effect of FY26 on predominantly liver PER2::LUC expression as a biomarker of the liver molecular circadian clock. The photon counts and locomotor activity records were stored on a connected PC. Data were then analysed with Spectrum Analysis (explained in “Statistical Methods”).
The RT-BIO enables bioluminescence recording from living organism mice with the aim to study circadian gene expression given the linkage of such gene to the luciferase sequence and the supply of luciferin in drinking water. Due to the expression of the gene luciferase fusion protein and its reaction with luciferin, photons can be detected by the PMT.

Two weeks after the start of entrainment, mice were implanted with the telemetric temperature sensor Anipill® as described in Section 2.8.2. Three days after surgery, mice were transferred into the RT-BIO for a 1-week adaptation including an additional synchronisation in the same LD12:12 as before surgery into this cage system. Mice were then exposed to constant darkness (DD). Feeding and drinking were *ad libitum*. With the intake of luciferin in drinking water, photons are produced as a by-product from the oxidation of luciferin to oxyluciferin and thus detected by the PMT. In order to minimise background noise related to the photons emitted by the bedding, mouse bedding was put into RT-BIO two days before the start of the experiment, so that such...
“noise” could be recorded. After 1 week in DD, mice were injected with either vehicle or FY26 (50 mg/kg i.p.) at ZT06 or at ZT18, then PER2::LUC expression was monitored in the RT-BIO for 7 additional days. The injection of mice required the removal of the mice from their cylindrical cages. This was done under dim red light. This possibly would have resulted in false positive photon counts detected by the PMT, if the PMT was not turned off and the shutter had not been closed before removing the cylindrical lead to take out the mouse for injection. Throughout the whole experimental duration, PER2::LUC expression and core body temperature were both recorded every 1 min and every 15 min, respectively. Data transfer between the Anipill® and the temperature receiver was accomplished on day 5 to 6 during treatment. Mice were visually checked every other day. The whole procedure was repeated for mice in Exp II and III (Figure II-16).

<table>
<thead>
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<th>Exp</th>
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<th>Mouse 2</th>
<th>Mouse 3</th>
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<tr>
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<td>ZT06: 50 mg/kg FY26</td>
<td>ZT18: 50 mg/kg FY26</td>
</tr>
<tr>
<td>II</td>
<td>ZT18: vehicle</td>
<td>ZT06: 50 mg/kg FY26</td>
<td>ZT18: 50 mg/kg FY26</td>
</tr>
<tr>
<td>III</td>
<td>ZT18: vehicle</td>
<td>ZT06: vehicle</td>
<td>ZT18: 50 mg/kg FY26</td>
</tr>
</tbody>
</table>

**Figure II-16**: Experimental set up of the effect of the PER2 expression in the liver clock.

(A) Three experiments were included in this study with one mouse entrained to LD12:12 and two other mice to antiphase LD12:12 cycles. (B) The entrainment of Exp II and Exp III started 2 weeks and 4 weeks later than Exp I onset.
2.11 Determining dose levels of FY26 to inhibit tumour growth

2.11.1 Aim

This study aimed to identify the therapeutic dose levels needed for FY26 to inhibit tumour growth. Such doses would then be used to determine whether tumour growth inhibition could differ according to circadian dosing time.

2.11.2 Experimental design

Tumour growth inhibition of FY26 was studied in 18 male CD1-Foxn1nu, aged 10 to 11 weeks, using the Hepa1-6 model. The mice were allocated to 4 cages (cages 1 to 4), harbouring 2, 3, and 4 mice each, respectively. Mice were kept in the common animal room of the BSU with lights on at 07:00 and lights off at 19:00. A cell suspension containing $1 \cdot 10^6$ Hepa1-6 Per2-luc cells in 100 µL PBS was prepared and kept on ice until injection. For each mouse, two syringes were loaded with 100 µL cell suspension. Mice under anaesthesia were injected subcutaneously (s.c.) with 100 µL cell suspension into the left and right dorsum. Tumour growth and body weights were measured for the subsequent 6 days. On day 7, then on every other day for 18 days, mice were injected with an intra peritoneal injection of vehicle, 40 mg/kg, 60 mg/kg or 80 mg/kg of FY26. Both, tumour inoculation and treatments, took place at clock hour 13:00, which corresponded to ZT06. Tumour was inoculated in the procedure room and drug was administered in the common animal facility. Throughout the experimental duration, body weight changes and tumour growth were monitored daily until 18 days after the start of treatment (Figure II-17).

![Experimental time line](image)

Figure II-17: Time line of tumour growth inhibition study.
2.12 Circadian tumour growth inhibition of FY26

2.12.1 Aim

The purpose of this experiment was to seek whether dosing FY26 near the times of best or worst tolerability would affect antitumour efficacy.

2.12.2 Experimental design

Antitumour efficacy was assessed following FY26 dosing at circadian time points ZT6 or ZT18 using tumour growth as an endpoint in 20 male CD1-\textit{Foxn1}\textsuperscript{nu} mice. Mice were 4 to 5 weeks old and allocated to 6 cages containing either 4 or 3 mice each. Mice were entrained with opposite LD12:12 cycles to enable treatment at ZT06 or ZT18 at clock hours 13:00 and 13:15 respectively. Ten mice were entrained with L on from 07:00 to 19:00, and another 10 mice with L on from 19:15 to 07:15 for 4 weeks, prior to tumour inoculation. Tumour inoculation took place between 13:30 and 15:00 clock hour under either blue or red light using anaesthesia. Seven days after tumour inoculation, treatment was started. Four mice were injected with vehicle and 6 mice with 50 mg/kg FY26 i.p. every other day at ZT06 or at ZT18 for 11 days (6 injections; cumulative dose, 300 mg/kg). Body weights and tumour growth were measured every day until the day after the 6\textsuperscript{th} injection, day 16. All the mice were euthanized seven days after the last treatment (end of experiment) or whenever human endpoints were exceeded (Figure II-18).
Figure II-18: Experimental design of time dependent tumour growth inhibition study.

(A) Schedule of light dark 12:12 entrainment. (B) Overview of the experimental time line.
3 Statistical Methods

3.1 Calculation of the population doubling time

The population doubling time describes the average time a cell population needs to double. In order to compute and calculate the PDT, cells were counted (described in chapter 2.3) every 24 h starting 24 h after cell seeding for up to 72 h. The PDT was calculated for individual cell counts every 24 h until the last day of cell counting by using the equation shown in Figure II-19. The PDT for the whole growth curve was obtained by calculating the average of the individual 24-h PDTs.

\[
t_{PDT} = \frac{h \cdot \ln 2}{\ln \left(\frac{C_2}{C_1}\right)}
\]

*Figure II-19: Equation to calculate the population doubling time (PDT).*

3.2 Time series Analysis of *in vitro* bioluminescence

The LumiCycle Analysis software Version 2.56 (Actimetrics, Wilmette, USA) was used to upload, to analyse and to export time series data into Excel (Microsoft Office Professional Plus 2013, Redmond, USA). In order to depict the time series in a graph, the background signal (photon counts of an empty spot in the turntable) was subtracted from the raw data. The transformed data were plotted in Graph Pad Prism 6. The two-sided running average was calculated by the average of the photon counts in a 24 h time span and was then subtracted from the centre photon counts in each 24 h time span. For example, if the first photon count would have taken place at 15:49 clock hour and the last at 15:39 the next day, the average of all the photon counts obtained between 15:49 and 15:39 (next day) would have been subtracted from the photon counts measured at 03:45. Plotting the time series of the raw data and the time series of the detrended data would thus leave out the first 12 h of data (Figure II-20).
Methods and Materials

As shown in Figure II-20, bioluminescent reporter time series were first detrended, so as to visualise sinusoidal oscillations and to compute the rhythm characteristics, including period, acrophase and amplitude. The period of an oscillation describes the interval of time between two peaks. The acrophase is the time at which the peak occurs in a periodic cycle, and the amplitude measures the difference between the maximum value at the acrophase and the mean of the oscillation, also called MESOR for midline-estimating statistic of rhythm (Figure II-23). The LumiCycle Analysis software was used to determine the period, acrophase and amplitude. Time series data were uploaded and imposed to a polynomial order of 5 to 7. The period, acrophase and amplitude were then calculated from 2.5 days after the start of synchronisation until the end of the experiment. The acrophase was normalised through the subtraction of the acrophase from the period and presented as acrophase relative to the period. The area under the photon counts curve (AUC) were further computed for specific time spans. The mean and the standard error of the mean (SEM) of the mentioned parameters were determined.

3.3 Analysis of real-time polymerase chain reaction data

The obtained C\textsubscript{T}-values (cycle threshold) were analysed using the Fluoro-Software R&D 1.3.22 (HAIN Technologies, Byfeet, UK). The C\textsubscript{T}-values were defined as the number of cycles needed until the fluorescence signal was above the fluorescence background signal. The C\textsubscript{T}-values were normalised to the baseline and the threshold was set at the start of the linear phase of the fluorescence curve. The obtained data

Figure II-20: Example of raw (panel A) and detrended data (panel B) of bioluminescence time series.
were exported to Excel and the quantification of the gene expression was analysed by using the $2^{\Delta\Delta Ct}$-method, also called “Livak Method”. The method normalised the expression of the interest gene against the reference gene. In a first step, the difference between the $C_t$-value of the interest gene control and the housekeeping gene ($36B4$) control [$\Delta C_t$ (control gene)] as well as the difference between the treated gene and the treated housekeeping gene [$\Delta C_t$ (treated gene)] was calculated. Next the difference between the difference of the control gene and treated gene was calculated [$\Delta\Delta C_t$ (control gene – treated gene)] and the logarithm of the $\Delta\Delta C_t$-value was taken to base 2 ($2^{\Delta\Delta Ct}$) (Livak & Schmittgen, 2001; Schmittgen & Livak, 2008). As $C_t$-value of each technical replicate of one biological triplicate was calculated, the median of the three technical replicates was picked and graphically plotted. The median $C_t$-value of each technical replicate was statistically analysed using either ANOVA or Cosinor Analysis.

3.4 Analysis of circadian rhythm dependent pharmacodynamics endpoints of FY26

The number of apoptotic cells and that of cells in the different cell cycle phases in each sample were counted using a BD LSR Fortessa flow cytometer (BD Sciences, Berkshire, UK) and analysed with the Flowing Software 2.5.1 (by Perttu Terho, University of Turku, Finland). The untreated controls were used to set the condition for the analysis. Therefore, the results of the non-treated samples (controls) were depicted in dot plot diagram using the Flowing Software showing all single measured cells represented by a dot in the graph. The viable cells were situated in the lower left quadrant, the apoptotic cells in the right upper quadrant and the necrotic cells in the lower right quadrant. The gates were set separating the different cell population from each other. This template was then transferred to the treated cells. The number of apoptotic, necrotic and living cells were further normalised to the total number of events which were predefined as 10 000 cell counts per sample. The normalised data were depicted in Graph Pad Prism 6 and statistical test were conducted.

The analysis of the cell cycle phases was done using the same software. FACS files were uploaded and the total number of events (10 000 cell counts) were presented in a histogram. Using the untreated control, the different cell cycle phases, $G_0/G_1$-phase,
S-phase and G2/M-phase, were selected. As for the analysis of the apoptotic cells, the same template was used to determine the number of events in each phase of the cell cycle on the treated controls. The number of cells in each cell cycle phase was further normalised to the number of events and depicted in Graph Pad Prism 6.

3.5 Analysis of core body temperature and PER2 liver expression time series

The time series of mouse core body temperature records obtained with the Anipill® and the time series of the PER2::LUC liver expression recorded with the RT-Bio were analysed using Spectrum Analysis. Spectrum Analysis is based on the decomposition of periodic and frequently measured time series into different oscillations (Rayner, 2001). As the core body temperature was measured every 5 to 15 min, and the PER2 expression every min, the hourly average of the core body temperature and the PER2 expression were calculated. The hourly data were used to apply Sampling-Resampling Spectrum Analysis. A 6-h sliding window of a 1-day window size was used for the core body temperature and a 2-h sliding window of a 1 day window size for the PER2 liver expression. The period, acrophase and amplitude with a 1st harmonic (24 h period) and a 2nd harmonic (12 h period) was calculated and plotted. The period, acrophase and amplitude of core body temperature and PER2::LUC expression were calculated over the 24 h before and the 24 h after vehicle or FY26 injection. The data were analysed using Spectrum Analysis. The script was written by Doctor Qi Huang (Costa et al., 2013; Komarzynski et al., 2018).

3.6 Analysis of tumour growth

The tumour burden was assessed by measuring the horizontal and vertical diameter (mm) of each tumour using callipers. The weight of each individual tumour in a given mouse was estimated by multiplying the smallest diameter by itself and then with the larger diameter. The product was then divided by two (Figure II-21). The obtained quotient represented the estimated tumour weight in mg, a standard procedure used for experimental chemotherapy studies (Tampellini et al., 1998).
The tumour weights, collected throughout the experimental duration, were normalised to the tumour weight before the start of treatment and plotted in Graph Pad Prism 6. The changes in tumour weights according to drug dose or injection time were assessed by calculating the AUC and the slope of the linear regression of the tumour growth curve from the start of treatment until the end of experiment.

3.7 Survival analysis

Kaplan-Meyer Analysis was used to compare survival curves and to determine statistically significant differences between groups. Therefore, the number of observed deaths over a defined time span was computed in a survival diagram in Graph Pad Prism 6. After the defined end of the experiment, log-rank test was applied to test the statistical difference of the occurrence of an earlier death in one group as compared to the other one. The log-rank (Mantel-Cox) test (Figure II-22) calculates the probability between the number of expected events and the number of observed events. The equation calculates a p-value. For a p-value of p<0.05 the number of events between two groups is statistically different whereas a p-value of p>0.05 indicates that the difference between the groups is insignificant in view of the power of the experiment, that depends upon sample size and number of events (Goel et al., 2010).
Methods and Materials

![Image](image176x629to461x766)

\[ p(\log \text{rank}) = \frac{(O_1 - E_1)^2}{E_1} + \frac{(O_2 - E_2)^2}{E_2} \]

- \(O_1 =\) observed events of group 1
- \(O_2 =\) observed events of group 2
- \(E_1 =\) expected events of group 1
- \(E_2 =\) expected events of group 2

Figure II-22: Equation of the log-rank test to calculate the statistically difference between the survival of predefined groups.

### 3.8 Cosinor Analysis

Data obtained from sampling over time, regardless of in vitro or in vivo data were analysed using Cosinor Analysis. Cosinor Analysis describes a mathematical model which is able to estimate the pattern of a curve while eliminating the noise coming from raw data. Data sampled as a function of time were fitted to a cosinor function using the least square method. The least square method describes the difference between the squares of the measurements and the squares of the estimated values according to the sampling time point. The mesor, acrophase and amplitude were computed in order to quantify the circadian characteristics of the data. The mesor is defined as the midline estimating statistic of rhythm, i.e. the mean of the fitted cosine function. The acrophase represents the time of the maximum in the fitted cosine function. The amplitude is the difference between the maximum of the cosine function and the mesor. These parameters are given with their statistical confidence limits (Figure II-23) (Cornelissen, 2014; Refinetti et al., 2007).
In this thesis, data were fitted against a 24-h and 12-h period, also called 24-h and 12-h harmonic, with at least 4 data point per sample and measurements taken every 6 hours. Cosinor Analysis was done using IBM SPSS Statistics 24 (IBM, New York, USA) (Cornelissen, 2014; Li et al., 2013).

3.9 Analysis of variance

One-way, two-way or three-way ANOVA’s with or without repetitive measurements were used as a statistical tests. ANOVA stands for analysis of variance and was used to test whether there was a significant difference between groups. One-way ANOVA tested differences in one independent variable between groups. Multiple-way ANOVA’s compared groups regarding two or three independent variables and their interaction terms. The null hypothesis meaning that the population mean were equal in groups for all factors and no interaction between any two or three factors, was rejected if the $p$-value was calculated as $p<0.05$. ANOVA was conducted either using Graph Pad Prism 6 for the IC50-studies or RStudio.Ink, (RStudio, Boston, USA) for the pharmacodynamics and in vivo studies and the analysis of the period, phase or amplitude of in vitro time series data. The significance levels were defined as: $p<0.01$ (*), $p<0.001$ (**), $p<0.001$ (**), $p<0.0001$ (***), $p<0.00001$ (****). The null hypothesis was rejected with $p$-values > 0.05.
4 List of Reagents

The following table summarised the reagents mentioned this chapter. It states from which company they were purchased as well as their LOT number.

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<tr>
<td><strong>Tris Base</strong></td>
<td>Fisher BioReagents</td>
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<td><strong>Trifluoroacetic acid</strong></td>
<td>Sigma-Aldrich</td>
<td>STBF6942V</td>
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<td><strong>TRIzol® Reagent</strong></td>
<td>Ambion by life technology</td>
<td>14174601</td>
</tr>
<tr>
<td><strong>Trypan Blue stain</strong></td>
<td>Invitrogen by Thermo Fisher Scientific</td>
<td>1856900</td>
</tr>
<tr>
<td><strong>Silicon Grease</strong></td>
<td>RS Components Ltd</td>
<td>494-124</td>
</tr>
<tr>
<td><strong>Sulforhodamine B</strong></td>
<td>Sigma Aldrich</td>
<td>MKBT0470V</td>
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<tr>
<td><strong>SuperScript II</strong></td>
<td>Invitrogen by Thermo Fisher Scientific</td>
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</tr>
</tbody>
</table>
III Results - Temperature Synchronisation
1 Preamble

Investigating the *in vitro* chronopharmacology of FY25 and FY26 requires adequate circadian synchronisation of the cell populations to be used in population assays. The experimental conditions should meet the following specifications: (i) to synchronise different types of cancer cell lines with a circadian period (*a priori* ranging from 18 to 30 hours), (ii) to allow for cells to remain viable for a minimum duration of 5 days, so as to enable drug effects to be measured over a sufficient time span, (iii) to keep cells synchronised with a consistent period, amplitude and acrophase throughout the experiment.

It is reported that dexamethasone shock was able to synchronise the circadian clocks of rat fibroblasts (Balsalobre *et al.*, 2000). Further, it was described by the same team that exogenous temperature cycles mimicking core body temperature could also synchronise circadian rhythms of cultured fibroblasts (Brown *et al.*, 2002). Dexamethasone shock was considered as a possible synchronisation method of murine hepatocarcinoma cells, using Hepa1-6 *Per2-luc* cell line as a model. Then, several candidate temperature cycles were tested to synchronise Hepa1-6 *Per2-luc* cells and to assess the consequence of genetic clock disruption on circadian entrainment through silencing clock gene *Bmal1* using the Hepa1-6 *Per2-luc shBmal1* cell clone. Furthermore, human colon cancer cell lines (HCT116 *Per2-luc* and HCT116 *Bmal1-luc*) were subjected to the established “optimal” temperature cycle in order to determine its synchronisation properties on different cancer cell lines. Cell culture conditions were further optimised according to initial cell counts seeded and to the luciferin concentration used. The synchronisation properties of the temperature schedules tested, were determined for the circadian clock using bioluminescence reporters and mRNA circadian expressions of core clock genes *Per2, Bmal1* and *Rev-erba*. Further, I have explored the relationship between the circadian clock and the cell cycle at a cell population level using temperature cycle synchronised Hepa1-6 cells with and without *Bmal1* silencing.
2 Synchronisation using dexamethasone shock

In a preliminary study, 5% or 10% FCS in cell culture media proved adequate for properly growing Hepa1-6 Per2-luc cells (not shown). Thus, 5% FCS was used in media concentration for all subsequent *in vitro* experiments. Luciferin was added to the media at a final concentration of 100 µM, except when mentioned otherwise.

To test whether a 30-min dexamethasone shock was a suitable method resulting in regular oscillations of Hepa1-6 cells, 75 000 Hepa1-6 Per2-luc cells were exposed to 1 mM dexamethasone for 30 min. *Per2-luc* bioluminescence oscillations of raw and detrended data (subtraction of the running average as described in chapter II.2) were observed for 2 days after the onset of dexamethasone shock.

![Graph](image)

*Figure III-1: Synchronisation of Hepa1-6 Per2-luc cells by dexamethasone shock.*

Seventy-five thousand Hepa1-6 Per2-luc cells were incubated with 1 mM dexamethasone for 30 min. The experiment was done in duplicate. Bioluminescence was recorded starting directly after washout. The media always contained 5% FCS. Time “0.0” refers to the end of dexamethasone exposure, when the media was replaced by dexamethasone-free media.

Both replicates exhibited periods of 20.4 h and 24.3 h, with corresponding acrophases occurring at 15.8 h and 17.4 h (modulo the respective period) and amplitudes of 514 and 388 photon counts per second. Yet, a steady decline in photon counts per second by 85.4% for replicate A and 57.8% for replicate B was observed from 0.9 days (replicate A) and 0.8 days (replicate B) until day 2. This decline in photon counts per
Results

second was coherent with a severe dampening of the oscillations thus likely reflecting a substantial amount of cell deaths (Figure III-1 and Figure III-2).

![Graphs showing period, amplitude, and acrophase](image)

**Figure III-2:** Period, amplitude and acrophase (modulo the corresponding period) of Hepa1-6 Per2-luc after dexamethasone shock.

These parameters were computed from time “0”, defined as time of serum shock end, using the detrended data time series of each replicate over 2 days.

3 **Circadian synchronisation of cell populations with exogenous temperature cycles**

3.1 **Synchronisation of a murine hepatocarcinoma cell line with temperature schedules A and B**

Exogenous 24-h periodic temperature cycles were tested with the aim to identify those which would meet our preset requirements. Schedules A and B involved two (schedule A) and one (schedule B) 24-h periodic temperature cycles before subjecting the cells to constant 37°C until 5 days after the start of temperature synchronisation (chapter
II.1). Both schedules were applied to 100,000 Hepa1-6 Per2-luc cells cultured in 5% FCS-containing media.

Figure III-3: Synchronisation of Hepa1-6 Per2-luc by temperature cycles A and B.

Panel A depicts the raw and detrended bioluminescence curves of Hepa1-6 Per2-luc subjected to temperature schedule A. Panel B shows the raw and detrended data of Hepa1-6 Per2-luc under the subjection of temperature schedule B.

The raw data of Hepa1-6 Per2-luc bioluminescence curves are characterised by a broad distribution. This can be explained by a decrease in luciferase expression due to possible cell death or insufficient synchronisation. The high variability between the replicates on schedule A precluded the computation of a mean circadian period of Per2-luc expression, as individual period length ranged from 19.4 h to 32.6 h in the 5 replicates. In contrast, the raw data of Hepa1-6 Per2-luc obtained under temperature schedule B, were characterised by a small variability resulting in a mean period (± SEM) of 20.2 ± 0.6 h (Figure III-3). Such periods represented the endogenous period of Hepa1-6 Per2-luc. The same variability was observed for the amplitudes.
Under temperature schedule A, the amplitude exhibited broad single values ranging from 457 photon counts per second to 1803 photon counts per second. However, due to the small variability achieved under temperature schedule B, the mean (± SEM) amplitude was 85 ± 32 photon counts per second. Considering the acrophases, a small variability between single values was obtained under both temperature schedules. The mean (± SEM) acrophase under temperature schedule A occurred at 21.7 ± 0.9 h and under temperature schedule B at 13.9 ± 0.9 h (Figure III-4).

A t-test compared the periods, acrophases and amplitudes of the cells exposed to either temperature cycles. No statistically significant difference was found for the period (p=0.089) as a result of large variability. The acrophases and the amplitudes differed with statistical significance with respective p-values of 0.0006 and 0.025. This showed that although both temperature cycles synchronised Hepa1-6 Per2-luc to similar periods, differences were seen regarding acrophases and amplitudes. However, a synchronisation to a 24 h period was not achieved which led to the investigation of the temperature cycles C and D.

![Periods, acrophases and amplitudes of Hepa1-6 Per2-luc cells synchronised to temperature cycle A and B.](image)

*Figure III-4: Periods, acrophases and amplitudes of Hepa1-6 Per2-luc cells synchronised to temperature cycle A and B.*
The parameters were taken after 2.5 days until 5.5 days after the start of synchronisation from 100 000 Hepa1-6 Per2-luc cells synchronised to temperature schedule A or B. A t-test revealed no significant difference between the periods ($p=0.089$) but between the acrophases ($p=0.0006$) and the amplitudes ($p=0.025$).

### 3.2 Synchronisation of murine hepatocarcinoma cell lines with temperature schedules C and D

Here, we replaced the release into constant temperature in schedules A and B, with alternating temperature plateaus at 37°C for 12 h and 36°C for 12 h in schedules C and D respectively. Under both temperature schedules, oscillations lasted until day 5, with obviously reduced variability between the single replicates generated under temperature cycle D as compared to schedule C (Figure III-5). The raw data presenting the photon counts per second showed that, under synchronisation on temperature schedule C, the highest photon counts per second ranged between 519 (day 3.3) and 1166 (day 3.2). These figures were twice as high as those found in cells synchronised with temperature cycle D, with highest photon counts per second ranging from 236 (day 3.4) to 530 (day 3.5).
Figure III-5: Synchronisation of 100 000 Hepa1-6 Per2-luc cells to temperature cycle C and D.

Both schedules synchronised Hepa1-6 Per2-luc to respective mean periods (± SEM) of 25.2 ± 1.1 h and 25 ± 0.8 h. The acrophases occurred at 22.4 ± 1.3 h on schedule C and at 23.6 ± 1.7 h on schedule D (modulo their respective periods) with much less variability between the individual acrophases of the replicates, as compared to schedule B.

The detrended data of cells synchronised with temperature cycle C showed a large variability of single amplitudes ranging from 266 to 686 photon counts per second with a mean (± SEM) of 468 ± 87 photon counts per second. The mean amplitude on temperature schedule C was thus 21-fold as large as that on schedule D (22 ± 4 photon counts per second).

A t-test was performed on the periods, acrophases and amplitudes. No statistically significant differences were detected in the bioluminescence periods and acrophases on both temperature schedule (p=0.929 and p=0.573 respectively), whereas a statistical difference was seen in the amplitudes (p=0.007) (Figure III-6).
To conclude, both temperature schedules synchronised Hepa1-6 Per2-luc cells, yet with higher variability in amplitudes and acrophases for culture dishes on temperature cycle C as compared to temperature cycle D.

![Graph showing periods, phases, and amplitudes after synchronisation with temperature cycles C and D.]

Figure III-6: Periods, phases and amplitudes of 100,000 Hepa1-6 Per2-luc cells after synchronisation with temperature cycle C and D.

The parameters were taken after 2.5 days until 5.5 days after the start of synchronisation from 100,000 Hepa1-6 Per2-luc cells synchronised to temperature schedule C and D. A t-test was conducted showing no differences between the temperature cycle C and D for the period (p=0.929) and the phase (p=0.453). However, a statistically significant difference was seen between the amplitudes of Hepa1-6 Per2-luc cells synchronised under temperature cycle C and D (p=0.0006).

3.3 Subjecting hepatocarcinoma cells to different τ-cycles matching temperature schedule D

Here, the temperature schedule D was further modified in order to test its synchronisation properties in Hepa1-6 Per2-luc clocks, near the known limits of circadian entrainment. Hepa1-6 Per2-luc cells were therefore synchronised to a period
of 20 h (T=20 h) or to a period of 26 h (T=26 h). I further examined whether starting cell seeding counts at 50 000 or 100 000 mattered regarding such entrainment (Figure III-7).

**Figure III-7:** Effects of a shortened and prolonged exogenous period on the endogenous period of Hepa1-6 Per2-luc with both tested τ-cycles of temperature according to modified schedule D.

Fifty-thousand (Left panels) or hundred-thousand cells (right panels) were seeded in 5 Petri dishes each then they were subjected to modified schedule D whose temperature cycle had a period of either T=20 h (schedule E, upper rows) or, T=26 h (schedule F, lower rows). Bioluminescence was recorded for 1 min every 10 min for up to 7 days.

Both T-cycles generated oscillations lasting until day 6 for an initial seeding cell count of 100 000 and until day 7 for 50 000 cells, this latter seeding count being also associated to a low variability between replicates.
whether the initial cell count was 50 000 or 100 000 cells. In contrast, the periods of *Per2-luc* oscillations on temperature schedule F (T=26 h), displayed average circadian periods of 24.9 ± 2.06 h or 30.4 ± 0.2 h for initial seeding counts of 50 000 or 100 000 cells. The periods obtained after synchronisation to a 20-h exogenous period showed no significant difference between 50 00 and 100 000 starting cell counts (*t*-test with *p*-value of 0.096). Entraining 50 000 and 100 000 Hepa1-6 *Per2-luc* cells to a 26-h exogenous period led to statistically significant different periods (*t*-test with *p*-value of 0.029). The comparison between the synchronisation of 50 000 starting cell counts with temperature cycle E (T=20 h) and F (T=26 h) did not result in any significant difference (*t*-test with *p*-value of 0.049). Comparing 100 000 starting cell counts with temperature cycle E and F, however, showed a significant difference between the periods according to the T-cycle (*t*-test with *p*-value of <0.0001) (Figure III-8 and Table III-1). The data suggested that a more consistent synchronisation to the imposed period was achieved by temperature schedule E (T=20 h) as compared to F (T=26 h). This was further confirmed through comparing their SEMs. Indeed, the imposed 20-h period of temperature schedule E (T=20 h) was close to the endogenous period of the Hepa1-6 *Per2-luc* cells, which could have facilitated the synchronisation, in contrast to the large variability in periods for the replicates submitted to the 26-h periodic temperature schedule F (T=26 h).

The phases relative to the period of 50 000 and 100 000 starting cell counts were similar in *Per2-luc* expression under synchronisation of the temperature cycle E (T=20 h). Subjecting the same amount of Hepa1-6 *Per2-luc* cells to the temperature cycle F (T=26 h) resulted in a phase delay of 6 h for 50 000 cells and of 8 h for 100 000 cells (Table III-1). The *t*-test was significant comparing temperature cycle E (T=20 h) and F (T=26 h) with a starting cell count of 100 000 cells (*p*=0.0001) and comparing 50 000 with 100 000 starting cell counts under the synchronisation with temperature cycle F (*p*=0.004) (Table III-1).
Table III-1: Level of significance between the T-cycle with a 20- and 26-h period and between 50 000 and 100 000 starting cell counts.

<table>
<thead>
<tr>
<th></th>
<th>t-test (50k cells)</th>
<th>t-test (100k cells)</th>
<th>t-test (T= 20 h)</th>
<th>t-test (T= 26 h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20-h vs. 26-h</td>
<td>p=0.049</td>
<td>p&lt;0.0001</td>
<td>p=0.096</td>
<td>p=0.029</td>
</tr>
<tr>
<td>20-h vs. 26-h</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>50k vs. 100k</td>
<td>p=0.096</td>
<td>p=0.0303</td>
<td>p=0.004</td>
<td></td>
</tr>
<tr>
<td>50k vs. 100k</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Period</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phase</td>
<td>p=0.494</td>
<td>p=0.188</td>
<td>p=0.012</td>
<td>p=0.0004</td>
</tr>
<tr>
<td>Amplitude</td>
<td></td>
<td></td>
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</tbody>
</table>

The amplitudes, achieved under the synchronisation of temperature cycle F, were for both starting cell counts, 50 000 and 100 000 cells, 4.3-fold higher as the amplitudes under the synchronisation with the temperature cycle E. Those differences were not significant according to the t-test (50 000 cells: p=0.0494, 100 000 cells: p=0.188) (Table III-1). Comparing the amplitudes from the initial cell counts showed a 1.4-fold higher amplitude for a starting cell count of 100 000 cells as compared to 50 000 cells (Figure III-8). This was observed for the temperature cycle E and F. The t-test (50 000 vs. 100 000 cells) showed statistically significant results with p=0.012 in temperature cycle E and p=0.0004 in temperature cycle F (Table III-1).

To conclude, temperature cycle E (T=20 h) synchronised 50 000 starting cell counts of Hepa1-6 Per2-luc cells to similar periods and acrophases, while larger variabilities in both parameters were found under temperature cycle F (T=26 h). This suggested that the 20-h period of temperature cycle E was close to the endogenous period of Hepa1-6 Per2-luc. Further, the results demonstrated that the entrainment of a lower starting cell count of 50 000 cells to a 20-h period resulted in less variability suggesting that the number of cells played a crucial role in the synchronisation of a cell population.
Results

Figure III-8: Periods, acrophases and amplitudes after synchronisation with temperature cycle E ($T=20\, h$) and F ($T=26\, h$).

The periods, acrophases and amplitudes were calculated for 50 000 and 100 000 Hepa1-6 Per2-luc cells taken from day 2.4 to 5.4 and from day 2.8 to 5.8 respectively.

3.4 Synchronisation of clock altered cells with temperature cycle D

The objective of this experiment was to investigate the synchronisation properties of temperature cycle D in clock-altered cells. To this end, the experiment was conducted using Hepa1-6 Per2-luc cells with silenced Bmal1 expression.

To verify the extent of Bmal1 silencing in Hepa1-6 Per2-luc shBmal1, the Bmal1 mRNA levels were compared between Hepa1-6 Per2-luc shBmal1 and Hepa1-6 Per2-luc taken from confluent 35 mm NUNC Petri dishes with an initial cell seeding number of 50 000 cells cultured at constant temperature at 37°C. A 64% inhibition in Bmal1 mRNA expression was found in the shBmal1 clone as compared to the original
Hepa1-6 Per2-luc clone (1.8 ± 0.7 vs 5.1 ± 1.4 relative gene expression) with a p-value of 0.112 from t-test and Wilcoxon test with p-value of 0.25 (Figure III-9).

Figure III-9: Expression of Bmal1 in Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 (n=3).

The clock alteration in Hepa1-6 Per2-luc shBmal1 was further investigated by subjecting 50 000 Hepa1-6 Per2-luc shBmal1 cells to temperature cycle B where cells were released at a constant temperature of 37°C, following a single 24-h temperature cycle with a 5°C magnitude. The raw data in Figure III-10 further revealed that even though the number of photon counts per second increased with a peak at day 4, no synchronisation to a stable period, phase and amplitude was achieved. A mean period (± SEM) was calculated as 28.9 ± 1.6 h, with a mean acrophase as 18.4 ± 1.0 h and a mean amplitude of 34 ± 18 photon counts per second. Figure III-10 further showed that no sustained oscillations were detected in such conditions for Hepa1-6 Per2-luc shBmal1, in sharp contrast with the Bmal1-proficient parent clone (Figure III-3).
Results

Figure III-10: Bioluminescence (non-subtracted and subtracted) and periods, acrophases and amplitudes of Hepa1-6 Per2-luc shBmal1 cells under synchronisation of temperature cycle B.

Fifty-thousand Hepa1-6 Per2-luc shBmal1 cells were seeded in a 35 mm NUNC Petri dish (n=4) and subjected to temperature cycle B while recording bioluminescence for 1 min every 10 min over an experimental duration of 5 days.

In a subsequent experiment, the synchronisation properties of temperature cycle D were assessed on Hepa1-6 Per2-luc and its altered-clock clone. Therefore, 50 000 Hepa1-6 Per2-luc and 50 000 Hepa1-6 Per2-luc shBmal1 were synchronised with temperature schedule D for up to 5 days (Figure III-11).

The raw data presented with a consistent increase of bioluminescence with a maximum ranging from 687 to 1724 photon counts per second from day 3.8 to 4.6. As expected, sustained and consistent oscillations were obtained from Hepa1-6 Per2-luc synchronisation whereas the oscillations of Hepa1-6 Per2-luc shBmal1 varied with a triangular shape pattern. The mean ± SEM period was 23.7 ± 0.4 h and 23.7 ± 0.2 h.
for the Bmal1-proficient and the Bmal1 silenced Hepa1-6 clones respectively ($p$-value of 0.889 from $t$-test).

The mean ± SEM acrophase for the clock deficient clone occurred at 18.4 ± 1.4 h, i.e. with a 4.5 h advance as compared to that at 22.9 ± 0.5 h in Hepa1-6 Per2-luc ($p$-value from $t$-test of 0.017). In contrast, no statistically significant difference in amplitudes were found according to shBmal1 silencing (27.3 ± 4.4 in wt vs. 36.8 ± 9.0 photon counts per second in shBmal1, with $p$-value from $t$-test of 0.369) (Figure III-12).

Figure III-11: Synchronisation of 50 000 starting cell counts of Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 by optimised temperature cycle D.

This experiment pointed out that the temperature cycle D was able to synchronise Hepa1-6 Per2-luc under the established conditions and to entrain the Bmal1-silenced Hepa1-6 Per2-luc clone. Considering the “triangular” shape of the shBmal1 oscillation peaking regularly at the end of the 36°C plateau, as well as the differences between the acrophases in both clones, it was concluded that schedule D induced pseudo-rhythmic bioluminescence patterns. The latter ones likely reflected changes in
temperature sensitivity of Hepa1-6 Per2-luc shBmal1, whose severe clock alteration was shown through both a 64% inhibition of Bmal1 mRNA and the lack of consistent circadian oscillations in Per2-luc following the release in constant temperature.

Figure III-12: Synchronisation properties of Hepa1-6 clones after temperature synchronisation with schedule D.

Periods, acrophases and amplitudes of Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 after synchronisation with the optimised temperature schedule D. T-test revealed a significant difference between the acrophase of Hepa1-6 Per2-luc vs. Hepa1-6 Per2-luc shBmal1 with $p=0.017$. No statistically significance was observed between the period ($p=0.889$) and the amplitude ($p=0.369$).

3.5 Clock genes expressions in temperature-synchronised Hepa1-6 Per2-luc cells

The highest expression of Per2 mRNA was observed at T12 with $0.18 \pm 0.03$ relative gene expression (RGE) (mean ± SEM) and the lowest one at T24 with $0.04 \pm 0.01$ RGE. The circadian expression of Bmal1 was in antiphase to the expression of Per2. Thus, the maximum of the Bmal1 expression occurred at T24 with $2.46 \pm 0.6$ RGE.
and the minimum at T08 with 0.46 ± 0.16 RGE. Rev-erba mRNA expression peaked at T16, i.e. 4 h after that of Per2 with 1.39 ± 0.3 RGE (Figure III-13).

Figure III-13: Per2, Bmal1 and Rev-erba expression in Hepa1-6 Per2-luc under temperature synchronisation using temperature cycle D.

The gene expression was normalised to the house keeping gene 36B4. The graphs depict the mean of the median (n=3) and SEM.

Cosinor Analysis with a 24-h period was conducted on the clock gene mRNA expressions of Per2, Bmal1 and Rev-erba in Hepa1-6 cells synchronised with temperature cycle D. Bmal1 and Rev-erba circadian expressions showed an additional significant 12-h harmonic rhythm (Table III-2).
Table III-2: Results from Cosinor Analysis of the circadian mRNA expression of clock genes Per2, Bmal1 and Rev-erbα in Hepa1-6 Per2-luc cells synchronised with temperature cycle D (mean ± SEM, n=3).

<table>
<thead>
<tr>
<th></th>
<th>24 h period</th>
<th>12 h period</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Amplitude (RGE)</td>
<td>Acrophase (h)</td>
</tr>
<tr>
<td>Per2</td>
<td>0.4 ± 1.2</td>
<td>13.0 ± 1.4</td>
</tr>
<tr>
<td>Bmal1</td>
<td>0.4 ± 1.2</td>
<td>0.6 ± 1.3</td>
</tr>
<tr>
<td>Rev-erbα</td>
<td>0.3 ± 0.1</td>
<td>17.7 ± 1.4</td>
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The acrophase in Per2 mRNA expression was compared to that in Per2-luc in Hepa1-6 Per2-luc cells synchronised with temperature schedule D using an initial cell counts of 50 000 cells. The mean ± SEM acrophase of the Per2-luc reporter construct was located at 22.9 ± 0.5 h (Figure III-12) and that of Per2 mRNA at 13.0 ± 1.4 h (Table III-2). Thus, both acrophases differed by 9.9 h. The acrophase of the Per2-luc reporter construct only described the acrophase of the luciferase expression whose transcription is linked to the promoter of Per2. Thus, the acrophase form the Per2-luc reporter construct does not represent the acrophase of Per2. This explains why the acrophase of bioluminescence curve differs from that of Per2 mRNA expression. Nevertheless, the reporter construct was adequate to determine the synchronisation properties of temperature cycles.

3.6 Circadian cell cycle phases of Hepa1-6 Per2-luc and its altered clock clone

A cell cycle analysis examined whether a circadian rhythm regulated cell cycle stage progression in Hepa1-6 Per2-luc or Hepa1-6 Per2-luc shBmal1 synchronised with the optimised temperature cycle D.

Therefore, fifty-thousand Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 cells in 35 mm NUNC Petri dishes were subjected to antiphase temperature cycles D starting at 37°C in incubator 1 and at 32°C in incubator 2 as described in chapter II.1. Another
50 000 Hepa1-6 Per2-luc cells were not subjected to temperature schedule D and served as an unsynchronised control. After 2.5 days, non-synchronised and synchronised cells (5 to 6 replicates per time point) were harvested every 4 h in a 12 h working schedule. Using antiphase synchronisation schedules ensured the sampling of cells of a 24 h experiment. Thus, the same clock hour referred to 2 circadian time points. Following the cell cycle analysis protocol (chapter II.1), the number of Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 cells in G0/G1-phase, S-phase or G2/M-phase were determined excluding apoptotic cells from the analysis. Across all 6 time points, the average number of apoptotic cells was 4.9% with a maximum (mean ± SEM) at T14 (9.3 ± 1.3%) and a minimum at T18 (2.4 ± 0.1%) (Data not shown). No statistically significant difference was observed (p-value from ANOVA with 0.425).

The non-synchronised cells were harvested at the same three real time points as the synchronised cells (Figure III-14) in a 12 h working schedule. The highest number of cells (mean ± SEM) in G0/G1-phase occurred at T06/T18 with 34.6 ± 1.0% and the lowest one at T02/T14 with 32.1 ± 0.5%. Regarding the S-phase, a maximum of cells was observed at T10/T22 with 34.5 ± 1.6% and the minimum at T02/T12 with 32.1 ± 1.2%. The G2/M-phase showed similar maxima and minima as compared to G0/G1-phase. At T06/T18 a maximum with 35.4 ± 1.4% of cells were detected in G2/M-phase. The minimum occurred at T02/T14 with 31.8 ± 1.7% cells. The data showed a consistent level of cells measured at different time points. ANOVA revealed no significant difference between the counts in specific cell cycle phases according to sampling time (p=0.148 for G0/G1-phase, p=0.463 for S-phase and p=0.507 for G2/M-phase).
Next, the circadian rhythm of the cell cycle phases was investigated in Hepa1-6 Per2-luc and in Hepa1-6 Per2-luc shBmal1 (Figure III-15). The highest number of cells (mean ± SEM) distributed in G0/G1-phase was observed at T14 for Hepa1-6 Per2-luc with 36.9 ± 1.1% and at T06 for Hepa1-6 Per2-luc shBmal1 with 31.7 ± 0.3%. In S-phase, both Hepa1-6 Per2-luc and its altered clock mutant showed the peak of cell accumulation at T18 to T22 with 39.3 ± 2.0% to 39.1 ± 2.6% for Hepa1-6 Per2-luc and at T22 with 46.4 ± 4.8% for Hepa1-6 Per2-luc shBmal1. Regarding the G2/M-phase the highest number of cells were obtained at T02 and T10 for Hepa1-6 Per2-luc with 33.4 ± 2.5% and 33.4 ± 2.3%. On the contrary, Hepa1-6 Per2-luc shBmal1 showed the highest number of cells at T22 with 39.2 ± 1.4%. Cosinor Analysis was conducted to investigate whether a circadian rhythm was present in the above stated cell cycle phases after temperature synchronisation (Table III-3).

The results of the Cosinor Analysis confirmed that under temperature synchronisation, a 24-h rhythm in Hepa1-6 Per2-luc was observed in S-phase with a mean ± SEM acrophase at 17.3 ± 1.2 h and with an amplitude of 0.1 ± 0.03% (p=0.006). For the
altered clock mutant, a 24-h rhythmicity was detected in G₀/G₁-phase with an acrophase of 10.9 ± 1.4 h and an amplitude of 0.17 ± 0.06% (p=0.035).

Even though previous experiments have proven the ability of the temperature cycle D to synchronise Hepa1-6 Per2-luc cells, a circadian pattern of the different cell cycle phase distribution was only detected for the S-phase in Hepa1-6 Per2-luc and G₀/G₁-phase for Hepa1-6 Per2-luc shBmal1. The results suggest that temperature synchronisation with a 24-h period did not synchronise the cell cycle of both Hepa1-6 clones although the clock gene expression were antiphase (chapter 3.5).

Figure III-15: Circadian cell cycle phases of Hepa1-6 Per2-luc (n=11) and Hepa1-6 Per2-luc shBmal1 (n=6) after synchronisation with temperature cycle D.
Table III-3: Results from Cosinor Analysis of the cell cycle phase distribution in 24-h temperature synchronised Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 (mean ±SEM, n=6).

<table>
<thead>
<tr>
<th>Period</th>
<th>Mesor (%)</th>
<th>Amplitude (%)</th>
<th>Acrophase (h)</th>
<th>p-Value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Synchronised Hepa1-6 Per2-luc</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G₀/G₁-phase</td>
<td>24-h</td>
<td>33.2 ± 0.9</td>
<td>-</td>
<td>0.204</td>
</tr>
<tr>
<td>S-phase</td>
<td>24-h</td>
<td>35.9 ± 0.8</td>
<td>3.7 ± 1.0</td>
<td>0.006</td>
</tr>
<tr>
<td>G₂/M-phase</td>
<td>24-h</td>
<td>30.8 ± 0.1</td>
<td>-</td>
<td>0.49</td>
</tr>
<tr>
<td><strong>Synchronised Hepa1-6 Per2-luc shBmal1</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G₀/G₁-phase</td>
<td>24-h</td>
<td>25.3 ± 1.1</td>
<td>4.3 ± 1.6</td>
<td>0.035</td>
</tr>
<tr>
<td>12-h</td>
<td>25.3 ± 1.1</td>
<td>7.2 ± 1.6</td>
<td>4.4 ± 0.4</td>
<td>0.0003</td>
</tr>
<tr>
<td>S-phase</td>
<td>24-h</td>
<td>40.9 ± 0.1</td>
<td>-</td>
<td>0.395</td>
</tr>
<tr>
<td>G₂/M-phase</td>
<td>24-h</td>
<td>33.8 ± 1.3</td>
<td>-</td>
<td>0.919</td>
</tr>
<tr>
<td>12-h</td>
<td>33.8 ± 1.3</td>
<td>4.9 ± 1.9</td>
<td>8.9 ± 0.7</td>
<td>0.044</td>
</tr>
</tbody>
</table>
4 The effect of different luciferin concentrations on period, phase and amplitude

It was recently reported that extracellular luciferin concentration could alter circadian period, phase and amplitude of mouse fibroblasts synchronised with a temperature schedule involving the alternation of a 12-h plateau at 32°C with 12-h plateau at 37°C for 5 days (Feeney et al., 2016). Based on this publication, the effect of different luciferin concentrations on the period, acrophase and amplitude of Per2-luc expression was investigated in Hepa1-6 cells, using temperature schedule D. So far, all experiments were performed using 100 µM luciferin. The purpose of this study was to identify and confirm the luciferin dose level that ensured a stable period, acrophase and amplitude as well as guaranteed enough luciferin supply for an experimental duration of 7 days.

Temperature schedule D was applied to 50 000 Hepa1-6 Per2-luc cells cultured in media containing 50 µM, 100 µM, 200 µM, 400 µM, 800 µM or 1.6 mM luciferin. Three replicates were prepared for each concentration. The occurrence of the maximum of the photon counts per second (Table III-4) pointed out that a higher luciferin concentration (400 µM to 1.6 mM) resulted in higher photon counts per second which occurred 0.8 days later as compared to the lower luciferin concentrations (50 µM to 200 µM).

*Table III-4: Overview of the occurrence of the maximum photon counts per second according to the different luciferin concentrations.*

<table>
<thead>
<tr>
<th>Luciferin concentration</th>
<th>Day of maximum photons counts per second</th>
<th>Maximum average photons counts per second (± SEM), n=3</th>
</tr>
</thead>
<tbody>
<tr>
<td>50 µM</td>
<td>3.8</td>
<td>1126 ± 153</td>
</tr>
<tr>
<td>100 µM</td>
<td>3.7</td>
<td>1572 ± 204</td>
</tr>
<tr>
<td>200 µM</td>
<td>3.8</td>
<td>1568 ± 138</td>
</tr>
<tr>
<td>400 µM</td>
<td>3.5</td>
<td>2134 ± 294</td>
</tr>
<tr>
<td>800 µM</td>
<td>4.5</td>
<td>2261 ± 316</td>
</tr>
<tr>
<td>1.6 mM</td>
<td>4.5</td>
<td>2561 ± 320</td>
</tr>
</tbody>
</table>
The analysis of the detrended data revealed a consistent period (mean ± SEM) ranging from 24.1 to 25.2 h, for up to 6 days, irrespective of luciferin concentration (Table III-4 and Figure III-17) (p-value from ANOVA with 0.347). This suggested that luciferin did not affect the period of the Hepa1-6 clock. However, luciferin concentration was responsible for differences in circadian amplitudes and acrophases.

![Oscillation of Hepa1-6 Per2-luc depending on extracellular luciferin concentration](image)

**Figure III-16:** Oscillations of 50,000 starting cell counts of Hepa1-6 Per2-luc depending on the extracellular luciferin concentrations.

*Both figures shown the average (± SEM) of three replicates.*

Considering the raw data in Figure III-16, no toxic effect of luciferin was apparent. Moreover, a dose-response relation was reflected by consistent increments in circadian amplitude, from 35 to 173 photon counts per second along with increasing luciferin concentrations (p-value from ANOVA with 0.006). The acrophase (*modulo* the respective period) was also delayed as a function of luciferin concentration, from 16.1 h to 22.4 h (p-value from ANOVA with <0.001) (Table III-5).

The amplitude and the acrophase were further plotted as dose response curves according to luciferin concentration (Figure III-18). The dose response curve of the amplitude was fitted to an exponential function. The inflection point of the fitted exponential function of the amplitude against the luciferin concentration occurred at a luciferin concentration of 110.9 µM. The dose response curve of the acrophase followed a linear function.
According to the Michaels-Menten-Kinetics, a higher concentration of extracellular luciferin resulted in a higher rate of oxidation reactions with luciferase. Put into context with this experiment, more oxidation reactions and thus a higher number of photons were detected under increasing luciferin concentrations. This was supported by the increased amplitudes obtained by increased luciferin concentrations.

Table III-5: Relevance of luciferin concentration for the period, acrophase and amplitude of the bioluminescence curve of Hepa1-6 Per2-luc cells (mean ± SEM, n=3).

<table>
<thead>
<tr>
<th>Luciferin concentration</th>
<th>50 µM</th>
<th>100 µM</th>
<th>200 µM</th>
<th>400 µM</th>
<th>800 µM</th>
<th>1.6 mM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Period (h)</td>
<td>24.7±0.1</td>
<td>25.2±0.7</td>
<td>24.6±0.03</td>
<td>24.1±0.2</td>
<td>24.4±0.1</td>
<td>24.4±0.3</td>
</tr>
<tr>
<td>Acrophase (h)</td>
<td>16.1±0.3</td>
<td>19.7±2.6</td>
<td>19.3±0.2</td>
<td>18.4±0.7</td>
<td>20.9±0.1</td>
<td>22.4±1.3</td>
</tr>
<tr>
<td>Amplitude (photon counts/min)</td>
<td>35.1±8.9</td>
<td>97.9±5.4</td>
<td>189.7±43.9</td>
<td>237.4±32.8</td>
<td>209.4±34.1</td>
<td>172.8±9.1</td>
</tr>
</tbody>
</table>
This experiment pointed out that luciferin did not exert any dose dependent effect on the clock period of Hepa1-6 Per2-luc cells. However, the acrophase occurred later and the amplitude increased along with increasing luciferin concentrations. At a luciferin concentration of 100 µM, the amplitudes were “high enough” to study the effect of drugs on the Per2-luc reporter in follow-up studies. Regarding the acrophase, Table III-5 shows that a 100 µM, 200 µM and 400 µM luciferin concentrations did not change the acrophase. Whereas the acrophase was advanced for 50 µM luciferin and delayed for 800 µM and 1.6 mM luciferin. Considering all the different factors, a luciferin concentration of 100 µM was confirmed as the best option.

5 Synchronisation properties of the optimised temperature cycle D in a human colon cancer cell line

Temperature schedule D was further applied to the human colon cancer cell line HCT116 Per2-luc in order to test whether this cell line could also be synchronised by the temperature cycle D. Further, the 24-h temperature cycle was switched over to constant 37°C after day 5. Releasing cells into constant condition aimed to investigate whether the oscillation of HCT116 Per2-luc would persist in the absence of time cues, thus enabling the determination of the endogenous period of this human cancer cell line.
The synchronisation of 50 000 starting cell counts HCT116 Per2-luc revealed consistent oscillations with a stable amplitude until day 5. Following the release at constant temperature, the regular oscillations persisted until day 9. The mean period (± SEM) was 23.6 ± 0.1 h under entrained conditions with a 1°C change between 37°C and 36°C and vice versa every 12 h from day 2.5 to day 5. The period increased to 24.9 ± 0.1 h following the release of the cells at constant temperature at 37°C from day 5 to day 9.5. Such period lengthening was statistically validated by t-test with a p-value of <0.001.

The mean ± SEM bioluminescence amplitude increased 3.4-fold from 38 ± 9 to 127 ± 19 photon counts per second from the synchronised conditions (day 2.5 to day 5.0) to constant temperature (day 5.0 to day 9.5) (p-value from t-test with 0.001) (Figure III-19).
The acrophases remained stable both during the temperature synchronisation (24.3 ± 0.3 h) and following the release of the cells at constant temperature (22.7 ± 4.5 h). However the periods in both conditions differed significantly (p-value from t-test of <0.001). The lack of time cues likely accounted for rather large variations in the acrophases of Per2-luc expression among the replicates, as indicated with SEM of 4.5 h, despite very similar periods. This observation supported the occurrence of desynchronisation, a phenomenon which could be avoided through maintaining temperature synchronisation throughout.

In a second experiment (Figure III-20) the synchronisation properties of temperature cycle D were compared between HCT116 Per2-luc and HCT116 Bmal1-luc. The aim was to identify the phase relations between Per2-luc and Bmal1-luc expressions, so as to further support the presence of a functional clock in HCT116, as well as to verify the synchronisation of the clock genes promoters of Per2 and Bmal1 by temperature schedule D.

HCT116 cells with the luciferase reporter Per2-luc and Bmal1-luc presented both with sustained oscillations and a trend for desynchronisation after the release in constant conditions on day 6. The mean ± SEM periods were calculated as 24.0 ± 0.2 h and 24.6 ± 0.1 h in these respective conditions (p-value from t-test of 0.019).
Figure III-20: Synchronisation of Hepa1-6 Per2-luc and Hepa1-6 Bmal1-luc to temperature cycle D and determining the periods, acrophases and amplitudes.

The mean ± SEM acrophase of Bmal1-luc expression occurred at 20.1 ± 0.2 h, thus occurring 2.5 h later than that in Per2-luc expression, which was located at 17.5 ± 0.5 h (p-value from t-test with 0.001).

The amplitude of the HCT116 Bmal1-luc (469 ± 50 photon counts per second) was 8.7-fold as high as compared to HCT116 Per2-luc (54 ± 7 photon counts per second) although the starting cell counts of both HCT116 clones were 50 000 (p-value from t-test with <0.0001). The reason for the difference between both amplitudes were traced back to the design of the reporter construct.

Synchronising HCT116 Per2-luc and Hepa1-6 Per2-luc cells to temperature cycle D resulted in consistent oscillations. Interestingly, the same schedule applied to the same initial cell count synchronised both Hepa1-6 Per2-luc and HCT116 Per2-luc cells to ~24 h, yet the mean ± SEM acrophase of photon counts per second occurred at 22.9 ± 0.5 h and at 17.5 ± 0.5 h, respectively (p-value from t-test with <0.0001). This 5.4 h phase difference further reflect cancer cell line specificities in clock properties. Indeed, the endogenous circadian period that was determined at a constant temperature of
37°C differed profoundly between both cell lines. Exposing 100 000 Hepa1-6 Per2-luc cells to temperature cycle B resulted in a mean ± SEM period of 20.2 ± 0.6 h under constant temperature (Figure III-4). Interestingly, exposing 50 000 Hepa1-6 Per2-luc cells to temperature cycle D with 20-h period, synchronised the cells to a mean ± SEM period of 20.2 ± 0.1 h. Those results might suggest that the endogenous period of Hepa1-6 Per2-luc was close to 20 hours, whereas the endogenous period of HCT116 was 24.6 h.
6 Postamble

The aim of the experiments reported here was to establish a synchronisation method which would allow synchronisation of cancer cell lines to a stable period, acrophase and amplitude and to ensure synchronisation duration for at least 5 days.

The synchronisation with temperature schedule D with alternating 12-h plateaus at 36 and 37°C, after an initial 24-h cycle ranging from 32 to 37°C, led to a stable period, acrophase and amplitude. T-cycle experiments of temperature schedule D revealed consistent synchronisation of Hepa1-6 Per2-luc cells with a 20-h temperature cycle, whilst somewhat variable synchronisation was observed under 26-h temperature cycles.

Temperature cycle D was further used to investigate its synchronisation properties following severe molecular clock alteration resulting from Bmal1 silencing. The subjection of Hepa1-6 Per2-luc shBmal1 cells to constant temperature showed no sustained oscillations. However, “pseudo-rhythms” were observed under the synchronisation of temperature cycle D. The human colon cancer cells HCT116 transfected with Per2-luc or Bmal1-luc were synchronised by temperature cycle D to sustained and antiphase oscillations.

Temperature synchronisation thus proved as a suitable method to synchronise cancer cell cultures. It was hence used in the subsequent experiments that investigated the in vitro chronopharmacology of FY26.
IV Results *In vitro* Chronotoxicity
1 Preamble

Temperature cycle D was used to synchronise cancer cell lines in order to investigate time dependent changes in the cellular toxicity of the novel osmium drug candidate FY26.

The aim was to establish a concentration scale of different FY26 concentrations which would allow to study the time dependent changes of the IC₅₀ (half maximal inhibitory concentration = IC₅₀) of FY26. This was used as an index of the cellular toxicity of FY26 in Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 and HCT116 Per2-luc cells. The required specifications for such a concentration scale were: (i) the achievement of extreme cell survival rates near 100% and 0% so as to calculate the IC₅₀ and (ii) the definition of a core concentration row with narrow variations of FY26 concentrations, in order to ensure the precise determination of the time dependent changes in IC₅₀.

Toward this goal, the above-mentioned cancer cell lines were synchronised using temperature cycle D. After 2.5 days of synchronisation, the cell lines were exposed to the selected FY26 concentration scales at circadian times T04, T10, T16 or T22. After an exposure time of 24 h, the FY26-containing media was removed and replaced by drug-free media. The total cellular protein content was measured 96 h after the start of drug exposure. The measured absorption of the media controls were used to normalise the absorption values from the drug exposed cells. The normalised values were expressed as percentages.

The time dependent changes in the IC₅₀ of FY26 were compared between Hepa1-6 Per2-luc, its clock-altered clone Hepa1-6 Per2-luc shBmal1, and the human colon cancer cell line HCT116 Per2-luc.
2 Time dependent cytotoxicity of FY26 on clock proficient and clock altered mouse liver cancer cells

2.1 Establishing the concentration scale to assess time dependent changes of FY26 cytotoxicity in the selected cancerous cell lines.

The *in vitro* chronopharmacology was assessed through testing different ranges of FY26 concentrations on the clock proficient liver cancer cell line Hepa1-6 *Per2-luc*, on its clock altered clone Hepa1-6 *Per2-luc shBmal1* and on the human colon cancer cell line HCT116 *Per2-luc*. All cell lines were exposed to the concentration rows shown in chapter II.1.

Concentration scale A was characterised by 1:10 dilutions of FY26, in order to identify the range where the IC$_{50}$ was situated. Based on this, concentration scales B, C and D were established, with closer values within the region of the IC$_{50}$. All of them led to a dose response curve with a maximum cell survival near 100% and a minimum near 0% as required to calculate the IC$_{50}$ (Appendix Figure XI-1 and Figure XI-3).

The Hepa1-6 *Per2-luc* cells were exposed to concentration scales A, B and C, with C representing the optimised scale. Its clock altered clone Hepa1-6 *Per2-luc shBmal1* was exposed to concentration scales A, B and D, with D representing the optimised scales. Mean IC$_{50} \pm$ SEM was 5.3 $\pm$ 0.4 µM for Hepa1-6 *Per2-luc* and 6.2 $\pm$ 0.4 µM for Hepa1-6 *Per2-luc shBmal1* using the respective optimised concentration scales for both cell lines (*p*-value from *t*-test with 0.139) (Figure IV-1).

Based on these findings, time dependent changes in IC$_{50}$ were investigated using concentration scales E and F for Hepa1-6 *Per2-luc* and F and G for Hepa1-6 *Per2-luc shBmal1*. All three scales shared the same core FY26 concentration sequence to ensure a precise detection of the time dependent changes of the IC$_{50}$. The maximum FY26 concentration differed amongst the scales E, F and G, being 16, 30 and 21 µM respectively (Appendix Figure XI-2).
Figure IV-1: IC$_{50}$ of FY26 for Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 after the exposure to the concentration scale C (conc. scale C) and D (conc. scale D) (n=12, with mean ± SEM).

Regarding the human colon cell line HCT116 Per2-luc, concentration scales H and I achieved maximum and minimum survival rates near 100% and 0%, thus met the requirements for IC$_{50}$ calculations (Figure IV-2 and Appendix Figure XI-3). In two separate experiments, 8000 HCT116 Per2-luc cells were exposed to scale H. Mean ± SEM IC$_{50}$ were 0.7 ± 0.0 µM and 1.5 ± 0.2 µM in the first and second experiment, respectively. In a third experiment, using concentration scale I, the mean ± SEM IC$_{50}$ was 4.9 ± 0.3 µM. ANOVA revealed significant differences in IC$_{50}$ values according to the concentration scale (p<0.0001). Tukey’s multiple comparison test indicated statistically significant differences between the IC$_{50}$ of each pair of experiments, thus highlighting the large inter-study variability that resulted from an extreme IC$_{50}$ value estimation, along the concentration scale tested (Figure IV-2). Concentration scale I was selected to test the time-dependencies in FY26 toxicity for HCT116 Per2-luc.
Results

Figure IV-2: Mean ± SEM IC$_{50}$ of FY26 using concentration scales H and I on HCT116 Per2-luc with an initial seeding count of 8,000 cells (n=4 to 8 with mean ±SEM).

3 Time dependent changes of the IC$_{50}$ of FY26 on Hepa1-6 Per2-luc cells

Time dependent changes in FY26 IC$_{50}$ were investigated in synchronised Hepa1-6 Per2-luc cells, within two experiments using concentrations scales E and F, both of which meeting the predefined specifications.

For both experiments, 4000 Hepa1-6 Per2-luc cells were initially seeded, and synchronised with temperature cycle D. After 2.5 days, cells were exposed for a fixed duration of 24 h to either concentration scales. Total cellular protein content was measured 72 h after FY26 withdrawal (Appendix Figure XI-4 and Figure XI-5).

Highly reproducible dose response curves were achieved in both experiments, involving 18 and 12 replicates respectively. Different IC$_{50}$ values were obtained according to the initial time of FY26 exposure. Mean ± SEM IC$_{50}$ values ranged from 4.1 ± 0.2 µM at T04 up to 5.9 ± 0.4 µM at T22 under concentration scale E, and from 9.6 ± 0.1 µM at T04 up to 12.3 ± 0.5 µM at T22 for concentration scale F. Thus, the IC$_{50}$ was consistently higher at T22 as compared to T04, by 44% and 28% under both respective concentration scales (Figure IV-3).
ANOVA showed a significant difference between time points for both experiments with a $p$-value of $p<0.0001$. Tukey’s multiple comparison test further emphasised statistically significant differences between T04 vs. T16 (***) and T22 (***), as well as between T10 vs. T22 (*) for the concentration scale E. The same test was conducted on the IC$_{50}$ values computed after exposing cells to concentration scale F. It showed a significant difference between T04 vs. T10 (*), T16 (*) and T22 (***) and between T10 vs. T16 (*) and T10 vs. T22 (*). Two-way ANOVA confirmed statistically significant differences between the IC$_{50}$ according to time ($p<0.0001$) and experiment ($p<0.0001$) without any interaction term ($p=0.219$).

**Figure IV-3: Circadian IC$_{50}$ of Hepa1-6 Per2-luc in two independent experiments.**

Panel A shows the dose response curve (left panel) and the IC$_{50}$ (right panel) according to time points of the concentration scale E ($n=18$ replicates) and panel B shows the dose response curve (left panel) and the IC$_{50}$ (right panel) after the exposure of the concentration scale F ($n=12$ replicates).
To conclude, the highest IC<sub>50</sub> was observed at T22, i.e. 2 hours before the temperature was scheduled to increase from 36°C to 37°C in the incubator. This time point was associated with reduced FY26 toxicity.

The interexperimental differences between the IC<sub>50</sub> values at the different circadian time points might have been caused because the experiments were conducted on different days as well by the possible use of different concentrations scales. The median concentration in either scales was 6 µM FY26. However, highest concentrations amongst the scales differed. Whereas the highest concentrations of concentration scale E constituted of 16 µM and 12 µM FY26, the maximum concentrations of concentration scale F were 30 µM and 16 µM FY26. Strikingly, the lowest cell survival was obtained under the exposure to concentration scale E. The variability in the measurement of the total protein content might have been a further contributing factor.

4 Time dependent changes of the IC<sub>50</sub> of FY26 on Hepa1-6 Per2-luc shBmal1 cells

Next, possible time dependent changes in the IC<sub>50</sub> were investigated in the clock-altered Hepa1-6 Per2-luc shBmal1 clone, using the same methodological approach as described for Hepa1-6 Per2-luc cells. However, concentration scales F and G were used here in two separate experiments, each one consisting of 12 replicates. Both concentration scales met the predefined specifications (Appendix Figure XI-6 and Figure XI-7).

The mean ± SEM IC<sub>50</sub> values obtained under the use of the concentrations scale F were 8.7 ± 0.2 µM both at T04 and at T16. At T10, the mean ± SEM IC<sub>50</sub> was calculated with 10.7 ± 0.3 and at T22 with 14.1 ± 0.4 µM. The exposure to the concentration scale G resulted in IC<sub>50</sub> values ranging from 4.9 ± 0.2 µM at T04, and 5.9 ± 0.1 µM at T16 up to 7.6 ± 0.2 µM at T10 and 7.4 ± 0.5 µM at T22. Thus, a bimodal pattern was identified using either concentration scales with the highest IC<sub>50</sub> corresponding to initial FY26 exposure starting at T10 (Figure IV-4).
Figure IV-4: Circadian IC$_{50}$ of Hepa-6 Per2-luc shBmal1 in two independent experiments.

Panel A shows the dose response curve (left panel) and the IC$_{50}$ (right panel) according to time point of two independent experiment using the concentration scales F and panel B shows the dose response curve (left panel) and the IC$_{50}$ (right panel) after the exposure to the concentration scale G at different time points. Each experiment consisted of 12 replicates ($n=12$ with mean ± SEM).

For both concentration scales, ANOVA reported a significant difference between the IC$_{50}$ of FY26 according to the time of administration. The $p$-value was $p<0.0001$. Tukey’s multiple comparison test highlighted a difference between T04 vs. T10 (**), T16 (*) and T22 (***) between T10 vs. T16 (***), as well as between T16 vs. T22 (**) for concentration scale G. A difference between time points of FY26 administration was seen for concentration scale F between T04 vs. T10 (**), T22 (**), between T10 vs. T16 (**), T22 (***) and between T16 vs. T22 (**). Two-way ANOVA detected a significant difference between time points ($p<0.0001$) and between both experiments ($p<0.0001$). Further, a significant interaction between time and either experiments was obtained ($p<0.0001$).
To conclude, FY26 showed the lowest toxicity in Hepa1-6 Per2-luc shBmal1 both at T10 and at T22, that respectively referred to 2 h before the temperature rose from 36°C to 37°C and decreased from 37°C to 36°C. Thus, the data differed from Hepa1-6 $\text{Per2-luc}$ as the IC$_{50}$ was higher at T10 and T22 whereas in Hepa1-6 $\text{Per2-luc}$ T22 was clearly identified at the time with least toxicity.

5 Time dependent changes of the cellular toxicity of FY26 on synchronised human colon cancer cells

Concentrations scale I was used for exposing synchronised HCT116 $\text{Per2-luc}$ cells at T04, T10, T16 or T22, after 2.5 days of synchronisation with temperature schedule D. Concentrations scale I was adequate, since minimum mean ± SEM survival ranged from 7.9 ± 1.7% to 11.7 ± 2.2% and maximum survival varied from 94.6 ± 1.7% to 104.7 ± 2.2% (Appendix Figure XI-8). Two experiments were performed.

The mean ± SEM IC$_{50}$ values ranged from 1.1 ± 0.1 µM for T04, to 3.3 ± 0.4 µM for T10, 2.6 ± 0.3 µM for T16 and 3.6 ± 0.2 µM for T22 in the first experiment. In the second one, the mean ± SEM IC$_{50}$ also followed a bimodal pattern, with values ranging from 5.5 ± 0.2 µM for T04, to 7.1 ± 0.2 µM for T10, 4.9 ± 0.1 µM for T16, and 7.1 ± 0.3 µM for T22. Thus, initial exposure of the cells at T10 or T22 was associated with least toxicity, by 22.7% to 45% as compared to T04 or T16 according to experiment. ANOVA on each individual experiment resulted in a significant difference between the IC$_{50}$ values according to time with a $p$-value of $p<0.001$. Tukey’s multiple comparison test further confirmed a difference between the time points T04 vs. T10 (***) and T22 (***) as well as between T16 vs. T22 (*) of the first experiment. A significant difference between the time points T04 vs. T10 (***) and T10 vs. T16 (***) and between T04 vs. T22 (**) was also seen in the second replication study, 2nd experiment (Figure IV-5). Two-way ANOVA confirmed statistically significant differences in IC$_{50}$ according to both FY26 timing ($p<0.001$) and experiment ($p<0.0001$). A significant interaction was observed between the time and experiment ($p<0.0001$) meaning that the effect of time on the IC$_{50}$ values is different from the effect of the experiment on the IC$_{50}$ values.
Figure IV-5: Circadian IC$_{50}$ of HCT116 Per2-luc after FY26 exposure in two separate experiments using concentration scale I.

Panel A shows the dose response curve (left panel) and the IC$_{50}$ values according to time (right panel) after the exposure of the concentration scale I and panel B shows the replication experiment (n=5-12 for panel A, n=13 for panel B).

6 Comparison of circadian toxicity patterns between clock proficient and clock altered cancer cell lines

To further test the consistency of the results amongst the individual experiments, the IC$_{50}$ values of FY26 were normalised to the average IC$_{50}$ obtained separately in each experiment for Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 and HCT116 Per2-luc. The maximum normalised mean ± SEM IC$_{50}$ for Hepa1-6 Per2-luc was 119.8 ± 4.7% for T22 and the lowest one 87.4 ± 2.9% for T04. The minimum normalised IC$_{50}$ was 99.5 ± 2.8% at T10 and 93.6 ± 2.9% at T16. ANOVA revealed a statistically significant difference between the normalised IC$_{50}$ values according to time of FY26 exposure.
Results

(p<0.0001). Tukey’s multiple comparison further revealed statistically significant differences between T22 and T04 (***) and T16 (****) (Figure IV-6).

*Figure IV-6: Comparison of the combined data of the normalised IC$_{50}$ of FY26 in Hepa1-6 Per2-luc (panel A) and in Hepa1-6 Per2-luc shBmal1 (panel B).

Panel A shows the IC$_{50}$ of Hepa1-6 Per2-luc (n=30) and panel B presents the IC$_{50}$ of Hepa1-6 Per2-luc shBmal1 (n=24).

Regarding Hepa1-6 Per2-luc shBmal1, the highest normalised mean ± SEM IC$_{50}$ was 126.7 ± 4.2% at T22 and the lowest of 79.0 ± 1.8% at T04. The average normalised IC$_{50}$ for T10 and T16 were 109.6 ± 2.8% and 87.1 ± 1.7% respectively. ANOVA revealed a statistically significant difference between the IC$_{50}$’s according to the time of drug administration (p<0.0001). Further, Tukey’s multiple comparison test emphasised a significant difference between T04 and T10 and T22 (**), between T10 vs. T16 and T22 (**), T14 vs. T22 (***) and T16 vs. T22 (****) (Figure IV-6). Cosinor Analysis showed an acrophase for a 24-h rhythm at T20.9 for Hepa1-6 Per2-luc whereas no 24-h and 12-h rhythm was detected for Hepa1-6 Per2-luc shBmal1 (Table IV-1).
Table IV-1: Cosinor Analysis of the normalised IC₅₀ of Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 and HCT116 Per2-luc (mean ± SEM).

<table>
<thead>
<tr>
<th></th>
<th>Period</th>
<th>Mesor (concentration in %)</th>
<th>Amplitude (concentration in %)</th>
<th>Acrophase (h)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hepa1-6 Per2-luc</td>
<td>24-h</td>
<td>100.1 ± 1.7</td>
<td>11.0 ± 2.4</td>
<td>20.9 ± 0.9</td>
<td>0.0001</td>
</tr>
<tr>
<td></td>
<td>12-h</td>
<td>19.0 ± 3.3</td>
<td>11.9 ± 0.0</td>
<td>&lt; 0.0001</td>
<td></td>
</tr>
<tr>
<td>Hepa1-6 Per2-luc shBmal1</td>
<td>24-h</td>
<td>99.6 ± 0.1</td>
<td>-</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>12-h</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td>HCT116 Per2-luc</td>
<td>24-h</td>
<td>98.9 ± 2.5</td>
<td>11.5 ± 3.7</td>
<td>19.9 ± 1.2</td>
<td>0.004</td>
</tr>
<tr>
<td></td>
<td>12-h</td>
<td>31.6 ± 3.9</td>
<td>8.2 ± 0.01</td>
<td>&lt; 0.0001</td>
<td></td>
</tr>
</tbody>
</table>

Thus, Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 exhibited different toxicity temporal patterns with the highest IC₅₀ at T22 identified for Hepa1-6 Per2-luc. Hepa1-6 Per2-luc shBmal1 on the contrary, showed a higher IC₅₀ at T10 and at T22 thus presenting a bimodal pattern.

7 Comparison of circadian toxicity between Hepa1-6 Per2-luc and HCT116 Per2-luc cell lines

In the last section, the IC₅₀’s between the murine hepatocarcinoma cell line Hepa1-6 Per2-luc and the human colon cancer cell line HCT116 Per2-luc were compared (Figure IV-7).
Figure IV-7: Comparison of the combined data of the normalised IC$_{50}$ of HCT116 Per2-luc (n=18-25) with Hepa1-6 Per2-luc (n=30).

Panel A presents the IC$_{50}$ of HCT116 Per2-luc and panel B shows the IC$_{50}$ of Hepa1-6 Per2-luc.

At T22, the human colon cancer cell line HCT116 Per2-luc exhibited on average the highest normalised mean ± SEM IC$_{50}$ with 127.1 ± 4.7% and at T04 and the lowest with 74.6 ± 5.9%. The IC$_{50}$ at T10 was 107.4 ± 5.5% and at T16 86.4 ± 3.8%. ANOVA confirmed a significant difference between the IC$_{50}$ values according to time with $p<0.0001$. Significant differences between T4 vs. T10 (***) and T22 (****) and T10 vs. T16 (*) and T22 (*) as well as between T16 and T22 (****) were shown after conducting Tukey’s multiple comparison test. Cosinor Analysis identified a 24-h rhythm with a mean ± SEM acrophase at T19.9 ± 1.2 h (Table IV-1). Compared with the highest and lowest toxicity of FY26 in Hepa1-6 Per2-luc, HCT116 Per2-luc needed 7.3% more FY26 at T22 to achieve a cell survival of 50% whereas at T04, 7.8% less FY26 was needed. This demonstrated that the time dependent IC$_{50}$ of HCT116 Per2-luc was characterised by a wider range as compared to Hepa1-6 Per2-luc. However, according to Cosinor Analysis, both cell lines were identified with the same time points of best FY26 tolerability. Two-way ANOVA showed a significant effect of FY26 timing on the IC$_{50}$ ($p<0.0001$) as well as a significant difference between both cell lines ($p=0.029$), with an interaction term ($p=0.028$).

The differences in the time dependent toxicity as well as in the magnitude of toxicity between Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 as well as HCT116 Per2-luc suggested that the different cell lines displayed different 24-h patterns in FY26 toxicity, which could be further altered according to the status of the molecular clock.
8 Postamble

The dose and dosing time dependencies in FY26 toxicity were determined using iterative IC_{50} determinations in the murine, clock-proficient hepatocarcinoma cell line Hepa1-6 Per2-luc and in the human colon cancer cell line HCT116 Per2-luc as well as in the clock-altered clone Hepa1-6 Per2-luc shBmal1.

In the absence of 24-h temperature synchronisation, the IC_{50} of FY26 was 20 to 30% lower in clock-proficient as compared to Bmal1-silenced Hepa1-6 Per2-luc cells, and 10% larger than that in human colon cancer cells HCT116 Per2-luc.

The synchronisation of cells with temperature cycle D revealed statistically significant 24-h changes in the IC_{50} of FY26 in both cancer cell lines and in the Bmal1-silenced Hepa1-6 Per2-luc clone. For Hepa1-6 Per2-luc, a circadian rhythm in FY26 toxicity was demonstrated. Best FY26 tolerability corresponded to a 24-h drug exposure starting at T22, 2 h before the programmed temperature rise from 36°C to 37°C. In contrast, both Hepa1-6 Per2-luc shBmal1 and HCT116 Per2-luc exhibited bimodal toxicity patterns with lowest toxicity at T10 and T22.

The next questions related to the ability of the in vitro chronotoxicity in 24-h synchronised Hepa1-6 cells to predict for in vivo chronotoxicity of FY26 in 24-h synchronised mice.
V Results – *In vivo* Chronotoxicity of FY25 and FY26
1 Preamble

The studies aimed to determine (i) the FY26 sublethal dose level to be used for subsequent circadian toxicity experiments, (ii) any circadian toxicity pattern (chronotoxicity) of FY26 and FY25 and (iii) organ toxicities caused by FY26. All experiments were conducted in mice synchronised with LD12:12.

The research questions were addressed in three different studies. The first study consisted of two experiments. Exp I and II tested 30, 40 and 60 mg/kg FY26 as potential sublethal doses. The second study involved three replicate experiments (Exp III, IV and V) for determining circadian toxicity pattern in FY26 (50 mg/kg i.p.), using body weight loss as main endpoint.

The effect of dosing time on body weight changes were analysed using three parameters: value at nadir, area under the curve over the 7 days following the day of injection (day 0), and the slope of recovery from nadir using linear regression. Overall survival analyses were also performed in order to further characterise the circadian toxicity patterns of FY25 and FY26.

Exp III to V not only served to determine the circadian toxicity patterns of both osmium complexes in mice, but also provided the bases for the subsequent studies investigating (i) FY26 effects on the circadian timing system of mice, using core body temperature and liver clock gene expression rhythms as circadian biomarkers, and (ii) tumour growth inhibition in hepatocarcinoma-bearing mice.

Exp VI involved the determination of histopathological lesions in the main organs of mice after subacute treatment with FY26.
2 Selection of sublethal dose levels of FY26 and FY25

For Exp I, 30 mg/kg and 40 mg/kg of FY26 were intraperitoneally injected in male C57BL/6 mice on day 1 and on day 5 respectively. The dose selection was based on a previous dose escalation study involving 20, 40, 80 and 160 mg/kg FY25 and FY26. The study identified 160 mg/kg as LD for FY26 (Appendix Figure XI-9). The injection of 30 mg/kg resulted in a maximum body weight loss of 7.2% for mouse 1 and of 8.1% for mouse 2 24 h after the first injection. No body weight loss was detected for the 3 other mice nor for the vehicle treated mouse. Body weight loss remained within the limits of the human endpoints.

![Body weight loss after a single i.p. injection of 30, 40 and 60 mg/kg FY26.](image)

*Figure V-1*: Body weight loss after a single i.p. injection of 30, 40 and 60 mg/kg FY26.

*Panel (A) shows the body weight loss after the treatment with 30 or 40 mg/kg FY26 and panel (B) shows the body weight loss after the treatment with 40 and 60 mg/kg FY26.*

Five days after the injection of 30 mg/kg FY26, the same mice were injected with a single i.p. dose of 40 mg/kg FY26. One day later, a maximum body weight loss of 7.3% for mouse 2, 3.8% for mouse 3 and 6.6% for mouse 5 were observed. Mouse 3 and mouse 5 recovered from the injection of 40 mg/kg FY26 and reached the pretreatment body weight 1 to 2 days after injection. Thus, mouse 2 lost 20% of body weight and exceeded the predefined human endpoints which led to its experimental exclusion. Throughout the whole experiment, the vehicle treated mouse did not lose any body weight (Figure V-1).

In Exp II, 3 mice were injected with a single intraperitoneal injection of 40 mg/kg and 60 mg/kg FY26. Maximum body weight loss was observed 24 h after drug injection
for the 3 mice treated with 40 mg/kg and for a single mouse of 3 mice dosed with 60 mg/kg. Two mice lost body weight down to 17.6% and 19.3% 3 days after treatment with 60 mg/kg FY26. Both mice were consequently euthanized for ethical reasons. Maximum relative body weight loss of the non-euthanized mice were 10.2%, 12.5%, 4.4% and 12.7% 24 h after the injection of FY26. All the non-euthanized mice recovered their respective pretreatment body weights 5 days after drug injection (Table V-1).

**Figure V-2: Survival of mice after treatment with 30 mg/kg, 40 mg/kg and 60 mg/kg FY26.**

*Panel (A) shows the survival after the treatment with 30 and 40 mg/kg FY26 and panel (B) shows the survival after the treatment with 40 and 60 mg/kg FY26.*

Mouse survival was plotted as a function of dose level according to Kaplan-Meier. Although no spontaneous death was encountered, the time of mouse sacrifice for ethical reasons (~20% body weight loss and/or poor appearance) was considered as the time of death. While dose levels 30 and 40 mg/kg were in the sublethal dose range, this was not the case for 60 mg/kg (Figure V-2). Nevertheless, the Log-rank test (Mantel-Cox) revealed no statistically significant difference in the survival of mice receiving 30 mg/kg or 40 mg/kg \( (p=0.368) \), or those treated with 40 or 60 mg/kg \( (p=0.114) \), possibly as a result of the limited sample size.

To summarise, the good tolerability of both lower doses tested, and the lack of statistically significant differences in both toxicity endpoints in these experiments, led to assume that a dose level of 50 mg/kg, i.e. midway between 40 and 60 mg/kg could ensure a decline in body weights which would be sufficient to detect circadian toxicity.
patterns without exceeding the predefined human endpoints. This dose level was chosen for the circadian testing of FY25 as well, since this compound presumably represents the active metabolite of FY26.

3 Circadian toxicity profile of FY25 and FY26

3.1 Distribution of mouse body weights before the start of treatment

The *in vivo* chronotoxicity was investigated in three consecutive experiments conducted on 234 male C57BL/6 mice aged 5 to 9 weeks. Body weights were monitored twice to three times a week during the pre-treatment synchronisation period from the day of arrival until the day of treatment.

The average mouse’s body weight (mean ± SEM) upon arrival for Exp III to Exp V was 18.5 ± 0.3 g for Exp III, 23.5 ± 0.4 g for Exp IV and 22.5 ± 0.5 g for Exp V. No differences in the body weights distribution amongst the groups was observed in any of the three experiments [p-value from ANOVA with p=0.747 (Exp III), p=0.575 (Exp IV) and p=0.913 (Exp V)]. On treatment day, the average mouse’s body weight was 27.9 g ± 0.2 g (Exp III), 27.9 g ± 0.5 g (Exp IV) and 25.6 g ± 0.6 g (Exp V). No significant differences of mouse’s body weight distribution amongst the injection time points were calculated for Exp III (p-value from one-way ANOVA 0.220) and Exp IV (p-value from ANOVA 0.939). Only in Exp IV, average mouse’s body weight was significantly different amongst the different injection groups (p-value from ANOVA 0.025). Thus, mouse body weights measured after the start of treatment were normalised to their pretreatment body weight (Figure V-3).

The analysis of the body weight distribution showed that no significant effect of time on the body weight development on untreated groups was observed except for the second experiment. The effect of 50 mg/kg FY25 and 50 mg/kg FY26 on the body weights according to the time of injection was thus studied on the normalised body weight data.
Figure V-3: Development of pretreatment mouse body weight distribution monitored from the day of arrival until the day of treatment.

3.2 Combined results of the circadian toxicity patterns of FY25 and FY26

3.2.1 Circadian changes of body weight loss after FY25 and FY26 administration

In order to assess the circadian toxicity pattern after a single intraperitoneal injection of 50 mg/kg of FY25 or FY26, the results from Exp III, IV and V were pooled, plotted and analysed together. As shown in Figure V-4, maximum normalised body weight loss occurred 24 h after FY25 or FY26 injection.

Mice receiving FY25, lost an average of 3.0 g (with average ± SEM body weight ranging from 26.6 ± 0.6 g to 23.6 ± 0.5 g) 24 hours after the treatment at ZT10. This corresponded to a relative decrease of mean ± SEM body weight of 10.9 ± 0.7%.
Average mouse’s body weight loss was smallest in mice receiving treatment at ZT10 as the average mouse boy weight dropped from 26.8 ± 0.6 g to 24.5 ± 0.5 g, corresponding to 8.3 ± 1.7% normalised body weight loss. The treatment of mice at ZT02, ZT06, ZT18 and ZT22 led to a decline of average mouse body weight of 9.1 ± 1.0%, 8.3 ± 1.3%, 9.4 ± 0.6% and 8.9 ± 0.9% respectively. Further, Figure V-4 showed that injected mice recovered and gained their pretreatment body weights, 5 to 8 days after the intraperitoneal injection of 50 mg/kg FY25 (Figure V-4).

Regarding FY26, maximum body weight loss was also seen 1 day after injection. Maximum weight loss was obtained at ZT18 and ZT22 with a decline from 100% pretreatment body weight of 9.0 ± 0.9% and 8.9 ± 0.9%. The minimum of body weight loss occurred when mice were injected at ZT02 (100% to 94.9 ± 1.1%). Following the curves of the body weight developments after injection of 50 mg/kg FY26, it was shown that mice reached their pretreatment body weight 4 to 8 days after drug administration (Figure V-4).

In order to assess the statistical significance of the dosing time related differences, two-way ANOVA with repetitive measurements was performed using the normalised body weight data, from the day of injection until the end of experiment for both FY25 and FY26. The Mauchly’s sphericity test demonstrated that FY25 and FY26 presented with a different toxicity profile (p<0.001) according to the time of drug administration (p<0.001). Further, ANOVA with repeated measurements was conducted on the body weight changes according to the time point of FY25 and FY26 injection resulting in a significant difference of body weight changes between time points after injecting 50 mg/kg FY25 (p<0.001) and 50 mg/kg FY26 (p<0.001). Tukey’s multiple comparison test showed a significant difference between the body weights at ZT2 vs. ZT14 (**) and ZT18 (*), ZT6 vs. ZT18 (**), ZT10 vs. ZT14 (*) and ZT14 vs. ZT18 (**) and ZT22 (*) after the treatment with 50 mg/kg FY25. Regarding the injection of FY26, a significant difference between body weights was observed between the time points ZT2 vs. ZT10 (*), ZT14 (**), ZT18 (**), ZT6 vs. ZT10 (*), ZT18 (**), ZT22 (*) and between ZT14 vs. ZT18 (*) and ZT22 (*).
Figure V-4: Average body weight changes as a function of the circadian timing of a single injection of 50 mg/kg of FY25 or FY26.

Panel (A) shows the body weight loss in g and panel (C) shows the normalised body weight loss after a single injection of 50 mg/kg FY25. Panel (B) and (D) presents the body weight loss in g and the normalised body weight loss after a single injection of 50 mg/kg FY26.

Next, ANOVA and Cosinor Analysis were conducted on the minimum body weight loss (nadir of body weight) following FY25 or FY26 administration. No statistically significant difference characterised maximal weight loss as a function of FY25 dosing time. However, maximum body weight loss differed significantly according to the time of injection of FY26 ($p=0.027$). Tukey’s multiple comparison test further showed statistically significant differences in maximum body weight loss following FY26 administrations at ZT2 vs. ZT18 (*) and at ZT2 vs. ZT22 (*) (Figure V-5).
Cosinor Analysis was applied to identify rhythmic patterns with 24-h or 12-h periods in relative maximum weight losses (Table V-1). No significant 24-h or 12-h rhythm was shown for FY25. In contrast, Cosinor Analysis revealed a significant 24 h rhythmicity in maximum body weight loss for FY26, with an acrophase at $6.7 \pm 3.3$ h and an amplitude of $1.7 \pm 1.3 \%$ ($p=0.039$).

Table V-1: Cosinor Analysis of nadir body weights, as percentages of pre-treatment values, after injection of 50 mg/kg FY25 or FY26 (mean $\pm$ SEM, n=18 per time point).

<table>
<thead>
<tr>
<th></th>
<th>FY25</th>
<th></th>
<th>FY26</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mesor (%)</td>
<td></td>
<td>Mesor (%)</td>
<td></td>
</tr>
<tr>
<td>Period</td>
<td>24 h</td>
<td>12 h</td>
<td>24 h</td>
<td>12 h</td>
</tr>
<tr>
<td>Acrophase (h)</td>
<td>-</td>
<td>-</td>
<td>6.7 ± 3.3</td>
<td>1.9 ± 1.9</td>
</tr>
<tr>
<td>Amplitude (%)</td>
<td>-</td>
<td>-</td>
<td>1.7 ± 1.3</td>
<td>1.6 ± 1.3</td>
</tr>
<tr>
<td>$p$-Value</td>
<td>0.564</td>
<td>0.498</td>
<td>0.039</td>
<td>0.061</td>
</tr>
</tbody>
</table>

Thus, the changes in body weight at nadir did not differ significantly in the mice receiving FY25. In contrast, a circadian toxicity pattern was demonstrated for FY26. ANOVA’s and Cosinor revealed that the lowest toxicity resulted from FY26 dosing during the early to middle of the light, i.e. during the first half of the rest phase of the rest-activity cycle. Highest toxicity was encountered in mice dosed in the second half of the dark span, i.e. during the mid- to late activity phase of their rest-activity cycle. Confirmation was further sought using the area under the curve (AUC), the slope of
body weight recovery and overall survival, as complementary chronotoxicity endpoints.

3.2.2 Circadian changes of the area under the curve of body weight curves

The areas under the normalised body weight curves (AUC\(_{0-7d}\)) were plotted against the time of injection (Figure V-6). The AUC\(_{0-7d}\) was used as a further measure for probing any circadian toxicity pattern, where the larger the AUC\(_{0-7d}\) value, the worse the toxicity.

![Figure V-6: Area under the curve (AUC\(_{0-7d}\)) from day of injection of 50 mg/kg FY25 (panel A) and FY26 (panel B) until 7 days after injection in % \cdot d.](image)

Regarding FY25, AUC\(_{0-7d}\) was least in the mice dosed at ZT18 (676.5 ± 1.8 %\cdot d) and highest in those treated at ZT14 with 684.9 ± 3.9 %\cdot d. However, no statistically significant difference according to FY25 timing was found according to both ANOVA (p=0.073) or Cosinor for periods of 24-h or 12-h (p=0.572 and p=0.204, respectively). In contrast, for FY26, the lowest AUC\(_{0-7d}\) was identified at ZT22 with 661.3 ± 12.6 %\cdot d and the highest one at ZT2 with 693.7 ± 2.7 %\cdot d. ANOVA demonstrated statistically significant difference between the AUC\(_{0-7d}\) according to treatment time with a p-value of 0.048. Tukey’s multiple comparison further emphasized significant differences between the AUC\(_{0-7d}\) of the time points ZT2 and ZT22 (*). No significant 24-h (p=0.091) and 12-h (p=0.089) rhythm was obtained for the AUC\(_{0-7d}\) after FY26 administration.

The analysis of the AUC\(_{0-7d}\) data confirmed previous results supporting the occurrence of a circadian toxicity pattern for FY26, with least toxicity occurring during the early
to middle of the light span, and highest toxicity occurring in the second half of the dark span.

3.2.3 Circadian recovery from the nadir of body weight loss to pretreatment body weights

Mice injected with 50 mg/kg FY25 exhibited a fast recovery at ZT14 with a mean reciprocal slope of 2.0 ± 0.2 body weight in %/h. The slowest or longest recovery was observed at the injection time point ZT18 with a reciprocal slope of 1.3 ± 0.3 body weight in %/h. ANOVA did not reveal any significant differences between the recovery after injecting 50 mg/kg FY25 ($p=0.271$). No significant 24-h or 12-h rhythm of the slope of recovery were detected (Cosinor Analysis: $p=0.747$) (Figure V-7).

In contrast, the fastest recovery after FY26 administration was seen at ZT6 with a slope of 2.4 ± 0.3 body weight in %/h and the slowest one being 1.3 ± 0.3 body weight in %/h at ZT18. These differences did not differ significantly using ANOVA ($p=0.203$) (Figure V-7). However, Cosinor analysis showed a significant 24-h rhythm with an acrophase at T 6.2 ± 1.5 h and an amplitude of 0.3 ± 0.1 body weight in %/h with a mesor of 1.7 ± 0.1 body weight in %/h ($p=0.034$).

![Figure V-7: Slope of linear regression from nadir of body weight loss to recovery.](image)

Panel (A) shows the slope of recovery after the treatment with 50 mg/kg FY25 and panel (B) presents the slope of recovery after the injection with 50 mg/kg FY26.

To summarise, recovery from maximum FY25 toxicity did not display any circadian rhythmicity, whereas this was the case for FY26, a result that was fully consistent with
those obtained using nadir or AUC$_{0-7d}$ normalised body weight values. ZT06 was identified as the time point of FY26 delivery that achieved best tolerability.

3.2.4 Circadian differences in overall survival after treatment with FY25 and FY26

The overall survival rates in the mice treated with FY25 or FY26 were 91.7 ± 2.8% and 88.9 ± 3.2% respectively. As a result, the overall survival curves of the mice in Exp III-V showed no statistically significant differences according to circadian timing using the Log-rank (Mantel-Cox) test ($p=0.189$ and $p=0.095$, respectively). Interestingly however, mortality was only observed following dosing of either compound at ZT14, Z18, or ZT22, when mice reached humane endpoints, thus confirming improved safety of both osmium compounds following dosing during the rest span of mice (Figure V-8).

![Graph showing overall survival of mice after treatment with FY25 and FY26.]

**Figure V-8**: Overall survival of mice after treatment with 50 mg/kg FY25 and FY26.

*Panel (A) shows the survival after the treatment with FY25 and panel (B) after the treatment with FY26.*

3.3 Histopathological lesions

Following the 4 repeated i.p. injections of either the vehicle or 50 mg/kg FY26 every other day in PER2::LUC mice, a maximum body weight loss after the vehicle injection ranged between 5.8% (32.3 g to 30.4 g) and 5.2% (30.8 g to 29.2 g) on day 4 following the second treatment. A maximum body weight loss ranged between 12.9% (30.3 g to
26.4 g) of mouse 3 on day 8 to 19.9% (33.0 g to 26.6 g) of mouse 6 on day 6. Mouse 4 was euthanized on day 3 following the clinical inspection (Figure V-9).

Figure V-9: Body weight changes after 4 repeated injections of the vehicle or 50 mg/kg FY26 in PER2::LUC mice.

The AUC from day 1 until day 3 (including mouse 3) was calculated (AUC\textsubscript{1d to 3d}). The AUC\textsubscript{1d to 3d} showed a statistically significant reduction of body weight with a mean ± SEM AUC\textsubscript{1d to 3d} for the vehicle of 199 ± 1 %·d and for the FY26 treated of 185 ± 1 %·d (\textit{p}-value from \textit{t}-test with 0.001).

Figure V-10: Area under the curve of body weight loss 24 h after the second treatment (AUC\textsubscript{1d to 3d}).
Tissue sampling was done 24 h after the last injection under consideration of the predefined human endpoints. Thus, only three mice were euthanized after completing the full treatment plan of 4 doses. One mouse was culled on day 3, and two other ones on day 6 or 7. Liver, both kidneys, spleen, sternum, stomach, ileum and colon were sampled, washed in PBS and fixed in 10% Neutral Buffered Formalin for 24 h before being transferred into PBS for tissue slicing.

No major toxicity was determined in stomach, ileum, colon, liver, kidney, spleen and bone marrow for both vehicle-treated mice. Mild steatosis in 10% of the hepatocytes was observed in the liver of a single FY26-treated mouse. Histopathological slides of the organs of the other two drugged mice showed no pathological changes (Appendix Figure XI-10, Figure XI-11, Figure XI-12, Figure XI-13, Figure XI-14, Figure XI-15 Figure XI-16).

4 Translating *in vitro* into *in vivo* chronotoxicity

The determination of the *in vitro* and *in vivo* FY26 chronotoxicity identified T22 and ZT06 as the time points of best FY26 tolerability. As the *in vitro* FY26 chronotoxicity was assessed by subjecting cells to temperature cycles mimicking the physiological core body temperature rhythm, the actual temperature cycle to which cells were exposed to in the cell culture incubator, was measured. This was done by placing a temperature sensor (Anipill®) in three 35 mm NUNC Petri dishes filled with cell culture media. In parallel, the Anipill® was implanted in the peritoneal of 3 mice in order to record core body temperature cycles. Cosinor Analysis was than applied on the hourly averaged *in vitro* and *in vivo* temperature cycles (Figure V-11 and Appendices Figure XI-17). The acrophase of the *in vitro* temperature cycle occurred at T6.7 ± 0.0 h with an amplitude of 0.6 ± 0.0°C and a mesor of 37.0 ± 0.0°C (p<0.001). The acrophase of the *in vivo* temperature cycle was in antiphase to the *in vitro* temperature cycle which occurred at T18.2 ± 1.0°C referring to the middle of the active phase. Its amplitude was identified with 0.6 ± 0.1°C and the mesor with 36.3 ± 0.1°C (p=0.001). Thus, the time point of best *in vitro* FY26 chronotoxicity, which was identified at T21, occurred 3 h after the bathyphase (nadir of temperature cycle). The same was observed for the *in vivo* FY26 chronotoxicity as the time point of best FY26 tolerability occurred during the bathyphase of the *in vivo* core body temperature cycle.
This leads to the conclusion that the lowest *in vivo* and *in vitro* toxicity occurred during the bathyphase of the temperature cycle.

![Graph showing temperature rhythm](image)

*Figure V-11*: In vitro (blue curve) and in vivo (red curve) core body temperature rhythm fitted to a cosine function.

*Cosinor Analysis was conducted on the in vitro temperature cycles (n=3) and on the mice core body temperature cycles (n=3). The marked lines depict the programmed temperature cycle and the white and grey background to the inactive and active phase of mice.*
5 Postamble

For FY25, no significant toxicity pattern regarding the changes in body weights, the AUC\(_{0-7d}\) as well as the survival was observed. On the contrary, FY26 toxicity differed significantly as a function of circadian timing, using body weight changes at nadir, as well as over 7 days (AUC\(_{0-7d}\)) or the slope for expressing recovery. The lowest and highest toxicities corresponded to FY26 dosing near the middle of the rest phase and near the end of the activity phase of the rest-activity circadian cycle of the mice, i.e. at ZT06 and ZT18 respectively. Cosinor Analysis further revealed that FY26 is best tolerated following its administration during the bathyphase of either temperature cycles.

The results clearly demonstrated that the toxicity profile of FY26 followed a circadian pattern, which is not the case for its active metabolite FY25.

Based on these findings, the following circadian studies aimed at the determination on the relationship between FY26 chronotoxicity and its effects on selected physiology and molecular circadian biomarkers. Specifically, this included the investigations of the circadian effect of FY26 on both core body temperature as a measurement of CTS disturbance and circadian clock gene \textit{Per2} expression in liver.
VI Results – *In vivo* efficacy of FY26
1 Preamble

The aim of this study was to determine whether (i) the FY26 activity that was demonstrated against Hepa1-6 *Per2-luc* cells *in vitro*, would be achieved *in vivo* and whether (ii) delivering FY26 at times of best or worst tolerability would impact on its efficacy in tumour-bearing mice.

In a first study, tumour growth rates were determined in 18 tumour-bearing male CD1-foxn1<sup>nu</sup> mice entrained to LD 12:12, with light onset at 07:00 and light offset at 19:00. Mice were injected s.c. with 10<sup>6</sup> Hepa1-6 *Per2-luc* cells in 100 μL PBS in each flank. One week after tumour cell implantation, mice were injected with 0, 40, 60 or 80 mg/kg of FY26 every other day for 12 days. Body and tumour weights were measured daily.

The second study aimed at assessing whether FY26 timing would differentially affect tumour growth inhibition according to whether a subacute treatment with FY26 was given at the time point of lowest (ZT06) or highest (ZT18) toxicity.
2 Determination of therapeutic dose levels of FY26

2.1 Body weight changes after repeated FY26 treatment

Mean body weights ± SEM were similar in the control and the three treatment groups on the first treatment day (controls: 30.3 ± 0.9 g; FY26 40 mg/kg: 30.6 ± 0.8 g; FY26 60 mg/kg: 30.5 ± 0.8 g; FY26 80 mg/kg: 30.3 ± 0.9 g; p-value from ANOVA with 0.992) (Figure VI-1). Thus, subsequent body weight changes could be ascribed to treatment effects.

![Figure VI-1: Body weight (g) distribution before the start of the six repeated injections with 40 mg/kg, 60 mg/kg and 80 mg/kg FY26.](image)

A dose dependent reduction of body weights was observed during treatment (Figure VI-2). Minimum average mouse’s body weight was measured 14 days after treatment onset, being 30.4 ± 0.3 g (94.0 ± 1.7% of body weight on the first treatment day) for the mice dosed with 40 mg/kg/injection, and with 28.4 ± 0.8 g (92.3 ± 3.0%) for those dosed with 60 mg/kg/injection. The largest average mouse’s body weight loss was observed 8 days after treatment with 80 mg/kg/injection when it reached 25.5 ± 0.3 g (87.3 ± 9.8%). Body weight loss did not differ significantly according the FY26 dosing (p-value from ANOVA with 0.407). A linear relation was found between FY26 dose and body weight (as a percentage of its pre-treatment value) in each mouse (Figure IV-2).
Results

Figure VI-2: Body weight changes following repeated treatment with 40 mg/kg, 60 mg/kg or 80 mg/kg of FY26 every other day for 12 days.

The left panel (A) shows the normalised body weights and the right panel (b) the lowest body weight reached in each mouse (as a percentage of its pretreatment value) as a function of FY26 dose.

To conclude, repeated treatments with 40 mg/kg or 60 mg/kg of FY26 induced maximum body weight loss of 6.0% and 7.7% three days after the 6th dose respectively. Subacute treatment with 80 mg/kg FY26 resulted in more than 20% body weight loss and was thus outside of the predefined humane endpoints. The body weight change dynamics on FY26 dosing documented here for CD1-\textit{Foxn1}nu differed from those shown in C57BL/6 despite same sex, similar age and same LD12:12 synchronisation (shown in III Result – \textit{In vivo} chronotoxicity of FY25 and FY26). More specifically, body weight dropped to a nadir 24 h post dose in C57BL/6 mice, but not in CD1-\textit{Foxn1}nu. These observations support strain-specific differences in FY26 toxicity profile.

2.2 Survival as a measure of antitumour efficacy

The survival rate varied from 0% in the mice receiving 80 mg/kg/injection to 40% for those treated with 60 mg/kg/injection and 62.5% for those receiving 40 mg/kg/inj. The survival rate in the control group was 33.3% because of tumour progression. Log-rank (Mantel-Cox) test revealed significant differences in survival curves with regard to the different treatment groups ($p=0.0001$) (Figure VI-3). The results highlighted a near doubling of survival in the mice dosed with 40 mg/kg/injection as compared to
controls, while treatment at higher dose levels did not improve survival or even were detrimental due to excessive toxicity.

![Graph](image1)

**Figure VI-3**: Survival of Hepa1-6 Per2-luc-12 days bearing mice receiving six alternate day injections of 40, 60 or 80 mg/kg/injection of FY26.

### 2.3 Tumour growth on subacute treatment with FY26

The distribution of tumour weights on the first treatment day revealed some imbalances between groups, with arithmetic mean ± SEM values ranging from 12.7 ± 1.5 mg for the vehicle group up to 40.8 ± 13.4 mg for the 80 mg/kg group. The differences were not statistically significant (p-value from ANOVA with 0.676). Nevertheless, tumour weight changes were compared as a function of drug dose using both raw data and percent changes relative to tumour weight on Day 1.

![Graph](image2)

**Figure VI-4**: Distribution of the tumour weight (mg) before the start of treatment with 40 mg/kg, 60 mg/kg and 80 mg/kg FY26.
In order to precisely determine any relation between tumour inhibition and FY26 dose, ANOVA with repeated measurements was applied to the normalised tumour weights over the initial 9 treatment days. Day 9 was chosen because all the mice treated with 80 mg/kg had to be culled for humane reasons before day 10. A non-statistically significant trend was shown with a $p$-value of 0.071.

Further, the area under the curve (AUC) from the tumour growth curve over time were calculated over both the initial 9 days when mice were alive in all groups ($\text{AUC}_{1\text{d to }9\text{d}}$) and over the whole Exp duration of 18 days ($\text{AUC}_{1\text{d to }18\text{d}}$) (Figure VI-5 and Figure VI-6).

*Figure VI-5: Tumour growth inhibition after six repeated injections with the mean ± SEM of vehicle, 40 mg/kg, 60 mg/kg and 80 mg/kg FY26.*

*Panel (A) shows the tumour weight in g and panel (B) shows the normalised tumour weights to the tumour weight upon the start of the treatment in percentage.*

Using $\text{AUC}_{1\text{d to }9\text{d}}$ as an endpoint, mean tumour growth inhibition was 3.3-fold for 80 mg/kg/injection, and 1.0- and 1.8-fold for 60 and 40 mg/kg/injection. $\text{AUC}_{1\text{d to }18\text{d}}$ further showed 2.0- and 1.5-fold tumour weight decrease on 60 and 40 mg/kg/injection doses levels as compared to controls (Table VI-1).
Results

Figure VI-6: Area under the curve of tumour growth inhibition from day 1 to day 9 (AUC\textsubscript{1d to 9d}) and from day 1 until the end of the experiment (day 18) (AUC\textsubscript{1d to 18d}).

Panel (A): median and range of AUC\textsubscript{1d to 9d} of tumour weights following FY26 treatments. Panel (B): median and range of AUC\textsubscript{1d to 18d} of tumour weights following FY26 treatments.

Non-parametric Kruskal-Wallis Test revealed statistically significant differences in both AUC\textsubscript{1d to 9d} ($p=0.02$) and AUC\textsubscript{1d to 18d} ($p=0.008$) (Figure VI-6). Post-hoc testing (Dunn’s Test) showed no difference between the treatment group for AUC\textsubscript{1d to 9d} but for AUC\textsubscript{1d to 18d} with a significant difference ($p=0.021$) was detected between the vehicle and 80 mg/kg FY26 group. As the survival of the 80 mg/kg treated group was 0% after 5 treatments, Kruskal-Wallis Test was conducted on the same date but leaving the 80 mg/kg treated mice out. The test demonstrated with a $p$-value of 0.327 that no difference in tumour growth inhibition of 40 and 60 mg/kg compared to the vehicle (control) group was detected.

Table VI-1: AUC\textsubscript{1d to 9d}, AUC\textsubscript{1d to 18d} and the slope of tumour weights after repetitive treatments with the vehicle, 40 mg/kg, 60 mg/kg and 80 mg/kg FY26 (mean ± SEM, $n=3$-5).

<table>
<thead>
<tr>
<th></th>
<th>vehicle</th>
<th>40 mg/kg FY26</th>
<th>60 mg/kg FY26</th>
<th>80 mg/kg FY26</th>
<th>Kruskal-Wallis/ANOVA</th>
</tr>
</thead>
<tbody>
<tr>
<td>AUC\textsubscript{1d to 9d} (tumour weight% ∙ time (d))</td>
<td>5778 ± 939</td>
<td>3120 ± 702</td>
<td>5389 ± 1282</td>
<td>1784 ± 382</td>
<td>$p=0.02$</td>
</tr>
</tbody>
</table>
### Results

| AUC_{1d to 18d} (tumour weight% \cdot time (d)) | 41051 ± 13981 | 20784 ± 7691 | 26789 ± 9719 | \( p = 0.327 \) |
| Slope (tumour weight/ time (d)) | 619 ± 110 | 251 ± 89 | 328 ± 91 | 137 ± 86 | \( p = 0.032 \) |

The slope of the linear regression adjusted to tumour growth allowed to differentiate slow (low slope) and fast (high slope) tumour growth, according to FY26 dose level (Figure VI-7). The maximum slope was found in the untreated control group.

Mice treated with 80 mg/kg showed a 4.5-fold decline in tumour growth slope. Treatment with 60 mg/kg and 40 mg/kg FY26 resulted in a 1.9- and 2.5-fold respective declines in tumour growth slope as compared to controls (ANOVA, \( p \)-value = 0.078).

**Figure VI-7**: **Slope of the linear regression of tumour growth under the treatment with 40 mg/kg, 60 mg/kg and 80 mg/kg FY26.**

To summarise, FY26 did exert a modest antitumour efficacy in mice bearing hepatocarcinoma tumour. FY26 efficacy was largest following dosing with 80 mg/kg/injection. However, this dose level was too toxic even though the drug was given at ZT06, i.e. the circadian time point of best FY26 tolerability. Minimal differences were found regarding the efficacy of 40 vs. 60 mg/kg/injection, with the latter schedule being more toxic than the former one. Based on these results, a dose of...
50 mg/kg/injection was chosen to further study time dependent effects of FY26 on
tumour growth inhibition in male CD1-\textit{Foxn1\textsuperscript{nu}} mice bearing Hepa1-6 \textit{Per2-luc}
tumours.

3 Relationship between chronotolerance and antitumour efficacy

3.1 Body weight loss after repeated FY26 injections at ZT06 vs. ZT18

On the injection day, mean ± SEM body weight was 30.5 ± 1.1 g in the vehicle group,
29.2 ± 0.8 g in the mice to receive FY26 at ZT06 and 29.2 ± 1.0 g for those to be
 treated at ZT18 (ANOVA, \( p = 0.536 \)). (Figure VI-8).

\[
\begin{align*}
\text{Vehicle} & : 30.5 ± 1.1 \\
\text{ZT06} & : 29.2 ± 0.8 \\
\text{ZT18} & : 29.2 ± 1.0
\end{align*}
\]

\[\text{ANOVA, } p = 0.536.\]

\textit{Figure VI-8: Body weight distribution before the start of 6 repeated treatments with 50 mg/kg FY26 every other day injected at ZT06 and ZT18.}

Figure VI-9 shows daily mean body weights expressed both as grams and as
percentages of pretreatment body weight. Control mice data were pooled whatever the
vehicle administration timing was. Mice on vehicle displayed a gradual increase in
body weight (Figure VI-9). In contrast mice on FY26 lost weight. Those treated at
ZT06, displayed a maximum body weight loss of 4.9 ± 1.2% on day 4 (after the 2\textsuperscript{nd}
dose). For those treated at ZT18, maximum body weight loss was 8.2 ± 1.6% on day
10, i.e. after the 5\textsuperscript{th} injection. ANOVA with repeated measurements showed a
statistically significant difference between tumour growth inhibition according to
FY26 dosing (non-normalised and normalised body weights with a \( p \)-value of
\(< 0.0001 \)). Significant differences were detected between the vehicle group vs. ZT06
Results

(****) and vehicle vs. ZT18 (****) for the normalised and non-normalised body weights. A significant difference was further detected between the normalised body weights of mice injected at ZT06 and ZT18 (**).

Figure VI-9: Development of body weight over time after the treatment with 6 repeated injections of 50 mg/kg FY26 at ZT06 and ZT18.

Panel (A) shows the body weight loss in g and panel (B) presents the normalised body weight loss in percentage.

3.2 Tumour growth inhibition after FY26 dosing at ZT06 and ZT18

Large differences in individual tumour weights were found before treatment onset, with extreme values ranging from 13.0 to 84.6 mg. The average ± SEM tumour weight ranged from 29.2 ± 0.8 mg up to 30.5 ± 1.1 mg according to mouse group, yet the differences were not statistically significant (ANOVA, p=.536) (Figure VI-10).
Results

Figure VI-10: Distribution of tumour weight (mg) before the start of treatments with 50 mg/kg FY26 at ZT06 and ZT18 or vehicle (control).

Tumour growth curves in the three groups are depicted in Figure VI-11. The right panel presents the non-normalised tumour weights (mg). The tumour weights were normalised to the mouse tumour weights before the start of treatment. Tumour growth was similar in controls and in mice receiving FY26 at ZT06. In contrast, tumour growth was delayed in the mice receiving FY26 at ZT18. ANOVA with repeated measurements showed a significant difference between the tumour weights in the vehicle, ZT06 and ZT18 group (non-normalised tumour weights (mg) with $p=0.005$ and the normalised tumour weights (%) with $p=0.009$). Post-hoc testing (Tukey’s multiple comparison test) revealed a significant difference between the tumour weights in mg of the vehicle vs. ZT06 group and between the normalised tumour weights (%) of the vehicle vs. ZT18.
Figure VI-11: Time dependent tumour growth inhibition after 6 repeated treatments with 50 mg/kg FY26 given at ZT06 and ZT18.

Panel (A) shows the tumour growth inhibition after the treatment with FY26 and the vehicle in mg. Panel (B) depicts the normalised tumour growth inhibition in percentage.

The slope of the linear regression representing the tumour growth inhibition was significantly different between treatment time points (p-value from ANOVA with 0.04). The slope from the linear regression of the tumour growth decreased 1.2-fold following the FY26 injection at ZT06. At ZT18, the tumour growth declined 2-fold as compared to the vehicle. Tukey’s multiple comparison test identified a statistically significant difference between the vehicle treatment vs. the treatment at ZT18 (*) (Figure VI-12 and Table VI-2).

Table VI-2: AUC<sub>1d to 16d</sub> and the slope representing the tumour growth after FY26 treatment at ZT06 and ZT18.

<table>
<thead>
<tr>
<th>Slope (mg/time(d))</th>
<th>Vehicle</th>
<th>ZT06</th>
<th>ZT18</th>
<th>One-way ANOVA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>253 ± 32</td>
<td>205 ± 41</td>
<td>131 ± 19</td>
<td>p=0.04</td>
</tr>
</tbody>
</table>

In summary, the experiment suggested maximum tumour growth inhibition following 6 repeated injections of 50 mg/kg FY26 every other day at ZT18, during the mid-span of the active phase of mice.
3.3 Survival of tumour-bearing mice treated with FY26 at ZT06 or ZT18

No death was recorded due to the vehicle toxicity. However due to fast tumour progression, mice died as a consequence of too high tumour weights on day 9 and 15. The overall survival of mice treated at ZT06 or ZT18 was 33.3%. Similar numbers of deaths were due to large tumour weights and toxicity. A trend toward least toxicity in ZT06-dosed mice was apparent. However, log-rank test did not reveal any statistically significant difference between treatment groups for overall survival ($p=0.481$) as well as for toxicity-related or tumour progression-related survival ($p=0.069$ and $p=0.592$ respectively).
Results

Figure VI-13: Time dependent survival of mice undergoing 6 repeated treatments every other day of either the vehicle or 50 mg/kg FY26 at ZT06 and ZT18.

Panel (A) depicts the overall survival whereas panel (B) depicts the number of death due to a high tumour weight and panel (C) shows the number of death due to high FY26 toxicity.
4 Postamble

The results presented in this chapter demonstrated that FY26 was modestly active against Hepa1-6 \textit{Per2-luc} cells at palpable stage. The efficacy depended upon dose level and it was limited by the toxicity of the largest dose level tested (80 mg/kg/d). FY26 seemed to be more active following dosing at the most toxic timing, a finding that would need confirmation on a larger number of mice. Subsequent studies aimed at determining the driver of the circadian toxicity profile of FY26.
VII Results – Time dependencies in pharmacokinetics and pharmacodynamics of FY26
1 Preamble

The experiments aimed to define both the circadian patterns in the in vitro cellular pharmacokinetics and pharmacodynamics of FY26, and the influence of the molecular circadian clock on FY26 pharmacodynamics.

The first pharmacokinetics experiment primarily aimed to determine the time dependent changes of intracellular FY26 uptake. Synchronised Hepa1-6 Per2-luc cells were exposed to 1 µM FY26 at six time points 4 h apart (T02, T06, T10, T14, T18 or T22) until the experiment was completed. Osmium concentrations were measured with Inductively Coupled Plasma Mass Spectrometry (ICP-MS) in extracellular and intracellular fractions collected over a time course of 72 h, representing a total of 360 [Os] determinations. Dosing-time dependent changes in FY26 pharmacokinetics and pharmacodynamics were assessed through the determination of maximum intracellular FY26 concentration ($C_{\text{max}}$), time to reach $C_{\text{max}}$ ($t_{\text{max}}$), area under the curve (AUC) and cellular viability in each culture dish (counts of dead and living cells per 1 mL cell suspension).

Three subsequent experiments aimed to determine the induction of apoptosis, autophagy and cell cycle phase distribution after FY26 exposure of Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 for 24 h (for apoptosis, autophagy and cell cycle phase distribution) and for 72 h (autophagy). A fifth experiment assessed possible time dependent cell cycle phase distributions after FY26 exposure. As the different pharmacodynamics endpoints were determined at different time points during the experiment, the effect of FY26 on bioluminescence was assessed under continues FY26 exposure which was monitored from the start of drug administration until 4 to 5 days after.
2 Time dependent cellular pharmacokinetic of FY26

2.1 Osmium concentration in Hepa1-6 Per2-luc cell culture media

Dosing-time dependent changes in FY26 pharmacokinetics were assessed in Hepa1-6 Per2-luc cells synchronised with temperature cycle D. After 2.5 days of synchronisation, 1 µM of FY26 was added into each of the 198 Petri dishes at circadian time points T02, T06, T10, T14, T18 or T22 (33 dishes per dosing time point) with T0 marking change from 36 to 37°C of the temperature cycle. The extracellular (media) and intracellular osmium concentrations were measured at 0 h (just before drug administration), then at 1 h, 3 h, 6 h, 9 h, 12 h, 18 h, 24 h, 30 h, 48 h and 72 h after drug administration (triplicate samples for each individual sampling time per circadian time). Osmium concentrations were determined using ICP-MS.

The extracellular osmium concentrations remained close to 1 µM, which was in accordance with the FY26 dose at all dosing times (Figure VII-1). However, a minimum of extracellular osmium concentration ($C_{\text{min}}$) was detected for each time point which varied across the different sampling time points. Table VII-1 summarises the distribution of the average $C_{\text{min}}$ that did not differ significantly according to FY26 timing. Declines in extracellular osmium concentrations likely resulted from FY26 cellular uptake. Throughout the whole experimental duration, the extracellular osmium concentrations ranged between 0.7 ± 0.1 µM and 1.2 ± 0.1 µM as a function of both dosing time and FY26 exposure duration. Overall, FY26 was abundantly supplied to ensure drug saturation in the media throughout the experiment.

Table VII-1: Extracellular osmium concentration (µM) of synchronised Hepa1-6 Per2-luc cells incubated with 1 µM FY26 (n=3).

<table>
<thead>
<tr>
<th>Circadian time</th>
<th>Time $C_{\text{min}}$ occurred</th>
<th>$C_{\text{min}}$ (osmium) in µM Mean ± SEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>T02</td>
<td>1 h</td>
<td>0.896 ± 0.036</td>
</tr>
<tr>
<td>T06</td>
<td>18 h</td>
<td>0.862 ± 0.022</td>
</tr>
<tr>
<td>T10</td>
<td>3 h</td>
<td>0.729 ± 0.141</td>
</tr>
<tr>
<td>T14</td>
<td>6 h</td>
<td>0.898 ± 0.045</td>
</tr>
<tr>
<td>T18</td>
<td>12 h</td>
<td>0.885 ± 0.01</td>
</tr>
</tbody>
</table>
No statistically significant difference between the minimum of the extracellular osmium concentration according to the time of the start of drug administration was detected (p-value from ANOVA with 0.352).

Figure VII-1: Osmium concentration of cell culture media subjected to Hepa1-6 Per2-luc cells.

The extracellular osmium concentration was measured at 0 h (time point of the administration of 1 µM FY26 thus serving as an untreated control), then at 1 h, 3 h, 9 h, 12 h, 18 h, 24 h,
30 h, 48 h and 72 h. The sampling time point “0 h” refers to the time when 1 μM FY26 was added. At the same time, media with drugs were taken and treated as blank controls. The osmium concentration in the media of each sample was determined using ICP-MS (n=3).

2.2 The cellular uptake of FY26 in Hepa1-6 Per2-luc is time dependent

The intra-cellular osmium concentration was determined according to the same experimental set up as used for the extra-cellular osmium determination. Figure VII-2 depicts the individual and averaged intracellular osmium concentrations over the 72 h following FY26 addition into the media at one of six circadian times.

![Graphs showing intracellular osmium concentrations over time.](image)

*Figure VII-2: Time dependent changes of intracellular osmium concentrations in synchronised Hepa1-6 Per2-luc (n=3).*
Based on the pharmacokinetics curves, the maximum of osmium intracellular concentration ($C_{\text{max}}$), the time needed to reach $C_{\text{max}}$ ($t_{\text{max}}$) and the AUC until $t_{\text{max}}$ ($\text{AUC}_{0-\text{t}_{\text{max}}}$) were determined. All three parameters were largest following FY26 addition at T02, and least following drug addition at T14 or T10. More specifically, $t_{\text{max}}$ (mean ± SEM) was 30.0 ± 0 h for the T02 group, as compared to 19 ± 8.2 h for T14. $C_{\text{max}}$ (mean ± SEM) was 28.7 ± 4.6 ng /10$^6$ cells for the T02 group, and 12.7 ± 2.1 ng/10$^6$ cells for the T14 group. $\text{AUC}_{0-\text{t}_{\text{max}}}$ (mean ± SEM) was 341 ± 8 ng · h/ 10$^6$ cells for the T02 group and 156 ± 14 ng · h/ 10$^6$ cells) for the T10 group (Figure VII-3).

**Figure VII-3: Time dependent intracellular osmium uptake.**

Panel (A) presents $C_{\text{max}}$, of intracellular osmium, panel (B) depicts $t_{\text{max}}$, and panel (C) shows the $\text{AUC}_{0-\text{t}_{\text{max}}}$ of the intracellular osmium uptake (mean ± SEM, n=3).

Cosinor Analysis validated statistically significant 24-h rhythms for both $C_{\text{max}}$ and $\text{AUC}_{0-\text{t}_{\text{max}}}$, for which a 12-h harmonic period was also found. No circadian rhythm was validated for $t_{\text{max}}$. The acrophases of $C_{\text{max}}$ and $\text{AUC}_{0-\text{t}_{\text{max}}}$ occurred both within the hour following temperature rise from 36 to 37°C, thus suggesting increased FY26 uptake following FY26 addition near this circadian time (Table VII-2). ANOVA confirmed statistically significant differences in $C_{\text{max}}$ values according to circadian
dosing time \((p=0.026)\). Tukey’s multiple test of comparisons emphasised on the difference between T02 vs. T14 (*). The AUC\(_{0h-t\text{max}}\) values also differed significantly \((p=0.006)\). Tukey’s multiple comparison test pointed out significant differences between T02 vs. T06 (*), T02 vs. T10 (**) and between T02 vs. T18 (*).

*Table VII-2: Results from Cosinor Analysis of FY26 cellular absorption (AUC\(_{0h-t\text{max}}\)) and maximum cellular concentration (C\(_{\text{max}}\)) according to drug administration timing.*

The results for \(t_{\text{max}}\) are not shown as they did not exhibit a 24-h or 12-h rhythm.

<table>
<thead>
<tr>
<th></th>
<th>24-h rhythm</th>
<th>12-h rhythm</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>AUC(_{0h-t\text{max}})</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Acrophase</strong></td>
<td>0.5 ± 1.0</td>
<td>2.4 ± 0.4</td>
</tr>
<tr>
<td>(time in h)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Amplitude</strong></td>
<td>0.2 ± 0.1</td>
<td>0.3 ± 0.1</td>
</tr>
<tr>
<td>(ng osmium per 10(^6) cells (\cdot) h)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>p-Value</strong></td>
<td>0.011</td>
<td>0.002</td>
</tr>
<tr>
<td><strong>C(_{\text{max}})</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Acrophase</strong></td>
<td>1.8 ± 0.9</td>
<td>-</td>
</tr>
<tr>
<td>(time in h)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Amplitude</strong></td>
<td>0.3 ± 0.1</td>
<td>-</td>
</tr>
<tr>
<td>(ng osmium per 10(^6) cells)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>p-value</strong></td>
<td>0.006</td>
<td>NS</td>
</tr>
</tbody>
</table>

In summary, a large amplitude circadian rhythm characterised FY26 uptake in temperature-synchronised Hepa1-6 *Per2-luc* cells. Drug uptake was about twice as large following drug addition at T02 as compared to T14 or T10, based on C\(_{\text{max}}\) or AUC\(_{0h-t\text{max}}\) values respectively. Such chronopharmacokinetics mechanism partly accounted for the *in vitro* chronotoxicity of FY26. Indeed, the maximum FY26 uptake and the highest toxicity resulted from FY26 addition at T02 and T04 respectively, i.e. during the early hours of the higher temperature plateau in the 24-h synchroniser. In contrast, there was no apparent relation between the times corresponding to lowest
Results

FY26 uptake (T10 or T14), and that associated with best tolerability, thus calling for further chronopharmacodynamics investigations.

3 Time dependent pharmacodynamics of FY26

The first experiment determined the effect of 1 µM FY26 on the time dependent cell viability changes. This was followed by two experiments which investigated the pharmacodynamics effects of FY26 on apoptosis and autophagy. Two further experiments determined the dose and time dependent effects of FY26 on cell cycle phase distribution.

3.1 FY26 displays time dependent inhibition of cell proliferation

Viable cell counts were determined iteratively over the 72-h that followed FY26 addition at one of 6 circadian times in synchronised Hepa1-6 Per2-luc cells. Cell counts increased steadily thereafter, yet with apparently different patterns according to dosing time (Figure VII-5). While an initial cell count of 50 000 cells had been seeded initially, the cell counts at 0 h (mean ± SEM), just prior to drug addition ranged, from $2.2 \times 10^5 \pm 3 \times 10^4$ to $4.3 \times 10^5 \pm 4.2 \times 10^4$ for a 1 mL cell suspension of an individual Petri dish. Statistically significant differences between the cell counts at different circadian time of sampling were detected ($p$ from ANOVA with 0.019) (Figure VII-4).

Figure VII-4: Number of cell counts of Hepa1-6 Per2-luc cells at the first sampling time point "0 h" at T02, T06, T10, T14, T18 and T22.

Tukey’s multiple comparison test further showed a significant difference between cell counts at T14 vs. T18 (*). Subsequent cell counts in each group were then normalised
to the average corresponding cell count at “0 h”. The AUC from 0 h to 72 h (AUC0h-72h) and the slope of the cell counts from the linear regression were determined. The results are shown in Table VII-3.

Figure VII-5: Normalised cell counts of Hepa1-6 Per2-luc cells.

The figures show the average ± SEM normalised (to the FY26 free control at 0 h) cell counts of Hepa1-6 Per2-luc cells. After 2.5 days under temperature synchronisation, 1 μM FY26 was added at 0 h referring to the circadian time T02, T06, T10, T14, T18 and T22. Cells were harvested and counted at each circadian time point 0 h, 3 h, 9 h, 12 h, 18 h, 24 h, 30 h, 48 h and 72 h. Cells per replicate of each condition (n=3) were counted.
The highest $\text{AUC}_{0h-72h}$ was observed at T10 and the minimum at T18. Cosinor Analysis found no significant 24-h or 12-h rhythm. ANOVA was significant with a $p$-value of 0.0003. Tukey’s multiple comparison test showed significant differences between the time points T02 vs. T06 (*), T10 (**), T18 (**), T22 (**) and T06 vs. T14 (*), T10 vs. T14 (**) and T14 vs. T22 (**) (Figure VII-6).

Table VII-3: Mean ± SEM of the $\text{AUC}_{0h-72h}$ and $\text{slope}_{0h-72h}$ of normalised cell counts according to the time of FY26 addition.

<table>
<thead>
<tr>
<th></th>
<th>T02</th>
<th>T06</th>
<th>T10</th>
<th>T14</th>
<th>T18</th>
<th>T22</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{AUC}_{0h-72h}$ (cell counts/ml ∙ h$^1$)</td>
<td>18806 ± 1906</td>
<td>21398 ± 1223</td>
<td>23352 ± 3311</td>
<td>19783 ± 4710</td>
<td>14012 ± 1768</td>
<td>21953 ± 2959</td>
</tr>
<tr>
<td>$\text{slope}_{0h-72h}$ (cell counts/h)</td>
<td>4.4 ± 0.5</td>
<td>5.0 ± 0.7</td>
<td>5.8 ± 0.8</td>
<td>5.2 ± 1.0</td>
<td>3.0 ± 0.5</td>
<td>3.2 ± 1.5</td>
</tr>
</tbody>
</table>

The maximum and minimum of the $\text{slope}_{0h-72h}$ were observed at T10 and T18. Cosinor Analysis revealed a non-significant trend with a $p$-value of 0.053 for a 24-h rhythm. ANOVA did not detect significant differences in $\text{slope}_{0h-72h}$ (Figure VII-6).

Figure VII-6: Time dependent changes of the $\text{AUC}_{0h-72h}$ (panel A) and the $\text{slope}_{0h-72h}$ (panel B) after the administration of 1 µM FY26.

Based on the cell counts, the 24 h doubling rate under continuous exposure of 1 µM FY26 was calculated (Figure VII-7). The fastest doubling rate (mean ± SEM) was observed at T10 and T14 with a doubling rate of $25.6 ± 2.2$ h and $25.5 ± 1.8$ h.
Following the drug administration at T18, cells needed on average (± SEM) 42.4 ± 1.9 h to double the number of cells. Thus, between T10/14 and T18 the doubling rate almost doubled. This was significant according to ANOVA with a $p$-value of $<0.0001$. Post-hoc testing (Tukey’s multiple comparison test) further showed significant differences between the doubling rate at the time points T02 vs. T18 (*), T06 vs. T18 (**), T10 vs. T18 (****), T14 vs. T18 (****) and T18 vs. T22 (**). Cosinor Analysis demonstrated a significant 24-h ($p=0.005$) and 12-h ($p<0.0001$) rhythm of the doubling rate of Hepa1-6 Per2-luc cells under continues FY26 exposure. The mesor was obtained with 30.7 ± 0.9 h. The amplitude and acrophase for the 24-h rhythm were 0.1 ± 0.01 doubling rate \cdot h and 20.9 ± 1.2 h. The amplitude and acrophase for the 12-h rhythm were accordingly calculated with 0.2 ± 0.03 doubling rate \cdot h and 5.6 ± 0.4 h.

Figure VII-7: Time dependent doubling rate of Hepa1-6 Per2-luc cells under the continues exposure of 1 \mu M FY26.

So far, least FY26 uptake was identified at T10 to T14. This is coherent with the doubling rate which was fastest at T10 to T14. However, cells proliferated fastest at T18 suggesting that under continuous FY26 exposure, the time point of best FY26 tolerability was identified at T18 (T21 according to Cosinor Analysis). This was consistent with the respective time points identified for the IC$_{50}$ as best FY26 tolerability was detected at T22. To better understand the link between cell cycle proliferation and the IC$_{50}$, the effect of FY26 on the cell cycle phase distribution was assessed next.
3.2 FY26 dose dependent effects on the cell cycle

Figure VII-8 depicts and Table VII-4 presents the results of the cell cycle phase distribution in Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 after the exposure to increasing FY26 concentrations.

The number of Hepa1-6 Per2-luc cells distributed in G0 and G1-phase dropped after the incubation of 5 µM FY26 and increased thereafter with increasing FY26 concentrations. Consistently, the cells in S-phase increased following the exposure of 5 µM and remained high after the incubation with 7.5 and 10 µM FY26. A decline of the number of Hepa1-6 Per2-luc cells in G2/M-phase was observed after the exposure of 7.5 and 10 µM FY26. This suggest that Hepa1-6 Per2-luc cells were arrested in S-phase and that cells continued to cycle from G2/M-phase to G0/G1-phase. ANOVA was significant for the distribution of cells amongst the cell cycle phases with \( p < 0.0001 \). No effect was observed for the FY26 concentration \( (p > 0.999) \) but for the cell cycle phase and FY26 interaction \( (p < 0.0001) \). Post-hoc testing (Tukey’s multiple comparison identified statistically significant differences between G0/G1- vs. S-phase (***) , G0/G1- vs. G2/M-phase (****) and between S- vs. G2/M-phase (****).

For the Bmal1-silenced cell clone Hepa1-6 Per2-luc shBmal1 the number of cells distributed in G0/G1-phase decreased after the incubation with 7.5 µM FY26. Consequently an increase of Hepa1-6 Per2-luc shBmal1 cells in S-phase was observed as well as a decrease of cells in G2/M-phase. Differently from Hepa1-6 Per2-luc cells, the Bmal1-silenced cell clones did not proceed to G0/G1-phase but entered apoptosis instead. ANOVA was significant for the distribution of cells in the cell cycle phases \( (p < 0.0001) \) but not the FY26 treatment \( (p > 0.999) \). The interaction between both factors was significant \( (p < 0.0001) \). Tukey’s multiple comparison test detected significant differences between G0/G1- vs. S-phase (***) , G0/G1- vs. G2/M-phase (****) and between S- vs. G2/M-phase (****).
Table VII-4: Dose dependent cell cycle phase distribution in Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1. Shown is the mean (± SEM) of three replicates (n=3).

<table>
<thead>
<tr>
<th></th>
<th>Hepa1-6 Per2-luc</th>
<th>Hepa1-6 Per2-luc shBmal1</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>G₀/G₁-Phase</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control (cell number %)</td>
<td>53.9 ± 1.1</td>
<td>55.3 ± 1.9</td>
</tr>
<tr>
<td>5 µM FY26 (cell number %)</td>
<td>49.2 ± 1.1</td>
<td>55.9 ± 0.5</td>
</tr>
<tr>
<td>7.5 µM FY26 (cell number %)</td>
<td>53.8 ± 1.2</td>
<td>52.8 ± 0.7</td>
</tr>
<tr>
<td>10 µM FY26 (cell number %)</td>
<td>56.7 ± 0.2</td>
<td>54.9 ± 1.1</td>
</tr>
<tr>
<td><strong>S-Phase</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control (cell number %)</td>
<td>13.8 ± 0.0</td>
<td>13.8 ± 1.0</td>
</tr>
<tr>
<td>5 µM FY26 (cell number %)</td>
<td>18.3 ± 0.6</td>
<td>10.9 ± 0.6</td>
</tr>
<tr>
<td>7.5 µM FY26 (cell number %)</td>
<td>16.5 ± 0.5</td>
<td>13.5 ± 0.2</td>
</tr>
<tr>
<td>10 µM FY26 (cell number %)</td>
<td>18.2 ± 0.9</td>
<td>20.6 ± 1.6</td>
</tr>
<tr>
<td><strong>G₂/M-Phase</strong></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure VII-8: Dose depending cell cycle phase distribution in Hepa1-6 Per2-luc (blue) and Hepa1-6 Per2-luc shBmal1 (red).
The number of apoptotic cells (Figure VII-9 and Table VII-5) was determined after the exposure of the same FY26 concentrations. Exposing both cell clones, Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1, to an FY26 concentration which was twice as high as the IC\textsubscript{50}, only resulted in 23.9 ± 1.4\% apoptotic cells for Hepa1-6 Per2-luc and in 35.5 ± 0.8\% apoptotic Hepa1-6 Per2-luc shBmal1 cells. Interestingly, a higher number of apoptotic cells were detected for the Bmal1-silenced Hepa1-6 Per2-luc cell clone. Incubating both clones with 7.5 µM FY26 doubled the number of apoptotic Hepa1-6 Per2-luc shBmal1 cells as compared to Hepa1-6 Per2-luc. These data are coherent with the decline of Hepa1-6 Per2-luc shBmal1 cells in G\_0/G\_1- and G\_2/M-phase confirming that a higher number of Hepa1-6 Per2-luc shBmal1 cells enter apoptosis instead of transitioning to G\_0/G\_1-phase \[ANOVA with \( p \text{-value of } p<0.001 \) for FY26 effect, with \( p=0.036 \) for the clock effect and \( p=0.001 \) for the interaction, Turkey’s multiple comparison test with significant differences between: controls of both Hepa1-6 clones (*), between the Hepa1-6 Per2-luc control and 10 µM FY26 of Hepa1-6 Per2-luc (**) and its clock altered clone (****), between 5 µM and 7.5 µM FY26 Hepa1-6 Per2-luc shBmal1 and 10 µM FY26 Hepa1-6 Per2-luc (**) and between 7.5 µM FY26 Hepa16 Per2-luc and 7.5 (*) and 10 µM FY26 (***), Hepa1-6 Per2-luc shBmal1 and between the control vs. 7.5 µM FY26 (***), 5 µM FY26 vs. 7.5 (**) and 10 µM FY26 (****) and between 7.5 µM and 10 µM FY26 (****)].
Results

Figure VII-9: FY26 dose dependent induction of apoptosis.

Table VII-5: Results of the normalised number (mean ± SEM of n=3) of viable and apoptotic cells according to the exposure of increasing FY26 concentrations.

<table>
<thead>
<tr>
<th></th>
<th>Hepa1-6 Per2-luc</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Vehicle</td>
<td>5 µM FY26</td>
<td>7.5 µM FY26</td>
<td>10 µM FY26</td>
</tr>
<tr>
<td>Viable cells (%)</td>
<td>88.5 ± 1.6</td>
<td>74.4 ± 7.9</td>
<td>78.1 ± 2.1</td>
<td>63.8 ± 1.4</td>
</tr>
<tr>
<td>Apoptotic cells (%)</td>
<td>7.2 ± 0.6</td>
<td>15.5 ± 5.4</td>
<td>13.5 ± 1.4</td>
<td>23.9 ± 1.4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Hepa1-6 Per2-luc shBmal1</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Vehicle</td>
<td>5 µM FY26</td>
<td>7.5 µM FY26</td>
<td>10 µM FY26</td>
</tr>
<tr>
<td>Viable cells (%)</td>
<td>86.1 ± 1.1</td>
<td>83.9 ± 0.8</td>
<td>52.8 ± 10.3</td>
<td>41.5 ± 1.4</td>
</tr>
<tr>
<td>Apoptotic cells (%)</td>
<td>4.6 ± 1.5</td>
<td>7.6 ± 1.0</td>
<td>30.0 ± 6.9</td>
<td>35.5 ± 0.8</td>
</tr>
</tbody>
</table>

As another cell death mechanism, the induction of autophagy under the exposure of 2.5 µM, 5 µM and 10 µM FY26 was investigated. The Figure VII-10 and Table VII-6 present the results of the LC3B mRNA expression immediately and 72 h after the end of the 24 h FY26 exposure in both Hepa1-6 Per2-luc clones (right panels). For both cell clones the LC3B expression was higher when measured immediately after the end of the 24 h drug exposure time as compared to a 72 h FY26 recovery time span. Moreover the LC3B expression showed a dose response with increasing LC3B levels.
following the incubation with increasing FY26 concentrations. Consistently with the apoptosis data, the LC3B levels in the Bmal1-silenced Hepa1-6 cell clone were higher immediately and 72 h after the end of the 24 h FY26 exposure. ANOVA did not reveal a significant effect of the clock ($p=0.856$) and time ($p=0.843$) on the LC3B expression, although a significant effect of the FY26 concentrations ($p=0.003$) was determined. Interactions were only significant for the FY26 concentrations and time with $p$-value of 0.016.

Figure VII-10: LC3B expression following the in exposure of 2.5 µM, 5 µM and 10 µM FY26 in Hepa1-6 Per2-luc (panel A) and Hepa1-6 Per2-luc shBmal1 (panel B).

Table VII-6: LC3B expression in Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 after the immediate end and 72 h after the exposure of 2.5 µM, 5 µM and 10 µM FY26 (mean ±SEM, $n=3$).

<table>
<thead>
<tr>
<th>Bmal1 silencing</th>
<th>RNA collection time</th>
<th>Relative gene expression (RGE) of LC3B</th>
<th>control</th>
<th>2.5 µM</th>
<th>5 µM</th>
<th>10 µM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No</td>
<td>24 h</td>
<td>0.3 ± 0.1</td>
<td>0.8 ± 0.5</td>
<td>0.5 ± 0.4</td>
<td>1.9 ± 0.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>72 h</td>
<td>0.8 ± 0.3</td>
<td>0.2 ± 0.1</td>
<td>0.6 ± 0.2</td>
<td>1.1 ± 0.3</td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>24 h</td>
<td>0.3 ± 0.2</td>
<td>1.5 ± 0.2</td>
<td>0.03 ± 0.02</td>
<td>0.9 ± 0.5</td>
<td></td>
</tr>
</tbody>
</table>
The data showed that FY26 led to a cell cycle arrest of Hepa1-6 *Per2-luc* and Hepa1-6 *Per2-luc shBmal1* in S-phase. Whereas Hepa1-6 *Per2-luc* cells continued to transition from G2/M-phase to G0/G1-phase, a higher number of Bmal1-silenced Hepa1-6 clone entered apoptosis pointing towards a clock dependent effect. Moreover, it was observed that the induction of autophagy was upregulated immediately after the end of the FY26 exposure. The *LC3B* levels decreased after a 72 h recovery time when the time dependent FY26 IC$_{50}$ was determined with the best tolerability at T22 based on the *in vitro* chronotoxicity.

### 3.3 Time dependent effect of FY26 on the cell cycle phase distribution

The time dependent effect of the cell cycle phase distribution after FY26 administration was tested at T04 and at T16 in both Hepa1-6 cell clones after the exposure of 7.5 µM FY26. This concentration was chosen as it was a compromising concentration between 5 µM and 10 µM FY26 which caused a higher number of Hepa1-6 *Per2-luc* and Hepa1-6 *Per2-luc shBmal1* cells distributed in G0/G1- and S-phase and a decline of cells in G2/M-phase.

A time dependent distribution of temperature synchronised Hepa1-6 *Per2-luc* cells was seen after the administration of the vehicle. From T04 to T16, the number of cells in G0/G1-phase decreased by 18.2% whereas the number of cells in S- and G2/M-phase increased by 5.2% and 13.2%. Thus, a shift of cells from G0/G1 to G2/M-phase was observed. Following the administration of 7.5 µM FY26 at T04, led to an increase of cells distributed in G2/M-phase of 10.8% whilst the number of cells in G0/G1-phase decreased by 10.8%. At T16 however, the number of cells in G0/G1-phase increased by 9.5% whereas the number of cells distributed in S- and G2/M-phase decreased by 2.2% and 7.3%. The results suggest that, a higher number of cells undergo apoptosis following FY26 administration at T04 as Hepa1-6 *Per2-luc* cells did not transition from G2/M-phase to G0/G1-phase. Thus, T04 constituted the time point at which a higher number of cells entered apoptosis under the exposure of 7.5 µM FY26. ANOVA showed that statistically significant differences between the distribution of
Results

cell amongst the different cell cycle phases ($p<0.0001$). Treatment ($p>0.999$) and time ($p>0.999$) did not affect the cell cycle phase distribution even though significant interactions were achieved between the cell cycle phases vs. time (***). 

No time dependent distribution of the temperature synchronised *Bmal1*-silenced Hepa1-6 cell clone after the vehicle administration was observed. At both time points the highest number of cells were distributed in G0/G1-phase. Following the incubation with 7.5 µM FY26 at T04 and T16, no time dependent difference in the distribution of cells amongst the cell cycle phases was observed. However, at both time points, the number of cells in G0/G1-phase decreased but increased in G2/M-phase pointing towards the induction of apoptosis regardless of the time of drug administration. ANOVA confirmed a significant difference between the cell cycle phase distribution ($p<0.001$). However, no significant effect of treatment ($p>0.999$) and time ($p>0.999$) was detected nor any significant interaction between the three factors.

![Figure VII-11: Time dependent effect of 7.5 µM FY26 on the cell cycle stages on synchronised and non-synchronised Hepa1-6 Per2-luc cells (n=3 for vehicle, n=4-5 for FY26).](image)
Table VII-7: Normalised number of cells in percentage (mean ± SEM) distributed in the cell cycle phases (n=3 for controls, n=6 for treated samples).

<table>
<thead>
<tr>
<th></th>
<th>Hepa1-6 Per2-luc</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>G0/G1-Phase</td>
<td>S-Phase</td>
<td>G2/M-Phase</td>
</tr>
<tr>
<td>Synchronised</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vehicle (cell number %)</td>
<td>T04</td>
<td>52.1 ± 0.9</td>
<td>17.9 ± 0.2</td>
</tr>
<tr>
<td></td>
<td>T16</td>
<td>33.8 ± 2.3</td>
<td>23.1 ± 1.2</td>
</tr>
<tr>
<td>Synchronised</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7.5 µM FY26 (cell number %)</td>
<td>T04</td>
<td>41.3 ± 5.7</td>
<td>17.9 ± 1.1</td>
</tr>
<tr>
<td></td>
<td>T16</td>
<td>43.3 ± 3.4</td>
<td>20.9 ± 1.5</td>
</tr>
</tbody>
</table>

|                       | Hepa1-6 Per2-luc shBmal1 |                     |                     |
|                       | G0/G1-Phase      | S-Phase             | G2/M-Phase          |
| Synchronised          |                  |                     |                     |
| Vehicle (cell number %) | T04  | 43.5 ± 1.0          | 30.3 ± 1.1          | 26.3 ± 0.1          |
|                       | T16  | 40.1 ± 0.3          | 30.4 ± 0.8          | 29.6 ± 1.0          |
| Synchronised          |                  |                     |                     |
| 7.5 µM FY26 (cell number %) | T04  | 34.7 ± 10.5         | 26.2 ± 2.2          | 39.2 ± 8.5          |
|                       | T16  | 36.9 ± 4.5          | 30.9 ± 2.9          | 32.2 ± 4.3          |

To conclude, the administration of FY26 caused an increase of apoptotic cells following the start of incubation at T04. This was not the case when FY26 was added at T16. Regarding the Bmal1-silenced Hepa1-6 cell clone, no time dependent effect was observed but therefore in increase of cells in G2/M-phase suggesting that cells enter apoptosis at T04 and at T16.

**4 Time dependent effect of FY26 on bioluminescence**

So far the in vitro chronotoxicity of FY26, as well as the effect of FY26 on cell growth, cell cycle phase distribution, induction of apoptosis and autophagy was determined. The determination of those endpoints took place either immediately or 72 h after the end of the 24 h FY26 incubation time. In order to combine this, the time dependent effect of FY26 on the bioluminescence was monitored 5 days after drug administration in Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 and in HCT116 Per2-luc cells.
4.1 Time dependent effects of 1 µM FY26 on Hepa1-6 Per2-luc bioluminescence

The specific circadian patterns in the cytotoxicity of a 24 h exposure to 1 µM FY26 in Hepa1-6 Per2-luc cells are recalled in Figure VII-12. At this dose level, the rate of viable cells (mean ± SEM) was 88.9 ± 2.3% following FY26 exposure at T04 as compared to 94.7 ± 1.4% at T16 (ANOVA, p=0.111, Cosinor Analysis p=0.054). Such cytotoxicity was in the range of an IC_{10} (inhibitory concentration which caused 10% cell death or 90% cell survival), with highest and lowest values matching previously-reported IC_{50} chronotoxicity data.

![Figure VII-12: Cell survival of Hepa1-6 Per2-luc after a 24-h exposure to 1 µM of FY26 (n=30).](image)

This dose level transiently inhibited Per2-luc bioluminescence, as quantified by the emission of photon counts per second. The exposure to media, vehicle or 0.2 µM FY26 did not alter the bioluminescence oscillation patterns (Figure VII-13). The bioluminescence curves of Hepa1-6 Per2-luc cells treated with media, vehicle and 0.2 µM FY26 at all four time points, reached the maxima on day 4 and declined until day 6. After day 6 a small increase in bioluminescence was observed. The largest and most sustained decline was observed after the administration of 1 µM of FY26 at T22 which lasted for up to 2 days. At T04, bioluminescence declined within 12 h then increased after day 4.
Results

**Figure VII-13**: Effect of FY26 on Per2-luc expression dynamics according to the time of drug administration.

The upper left panel shows the bioluminescence of the Per2-luc reporter in Hepa1-6 Per2-luc followed the administration of 1 µM FY26. The time point of drug administration was marked with a red hatched, vertical line. The upper right panels shows the bioluminescence of the same cell line followed FY26 exposure at T10, the lower left at T16 and the lower right at T22.

In order to analyse the time dependent differences of immediate FY26 effects on Per2-luc bioluminescence, the differences in the area under the bioluminescence curves were calculated from the time of drug administration until the bioluminescence curve of 1 µM of FY26 crossed the mean of the vehicle bioluminescence curves. The difference between the AUC_{vehicle} and the AUC_{1µM} was calculated and plotted as AUC_{Diff} (Figure VII-14). Maximum AUC_{Diff} occurred at T22 (1236 ± 244 photon counts per second · h). It was 3-fold as high as the smallest AUC_{Diff} that followed FY26 addition at T04 (414 ± 43 photon counts per second · h). ANOVA was significant with a p-value of 0.047 with a significant difference between T04 vs. T22.
Results

(*) according to Tukey’s multiple comparison test. Cosinor Analysis demonstrated no significant 24-h rhythm \((p=0.076)\).

**Figure VII-14:** Time dependent changes of the AUC\textsubscript{Diff} after the administration of 1 \(\mu\text{M}\) FY26 at T04, T10, T16 and T22.

To summarise, AUC\textsubscript{Diff} represents the FY26 effect on \(\text{Per2-luc}\) bioluminescence immediately after the drug administration. The highest inhibition of bioluminescence was observed following the drug administration at T22, as the difference between the AUC\textsubscript{Vehicle} and AUC\textsubscript{1\(\mu\)MFY26} was highest. The lowest bioluminescence inhibition occurred at T04. Thus, both time points showed opposite toxicity pattern as compared to previously demonstrated IC\textsubscript{50} chronotoxicity pattern (T22 best tolerated time point).

### 4.2 Time dependent effect of 1 \(\mu\text{M}\) FY26 on the \(\text{Per2-luc}\) reporter construct in Hepa1-6 \(\text{Per2-luc shBmal1}\)

A similar experimental set up comparing a single dose of 1 \(\mu\text{M}\) FY26 to vehicle was applied to \(\text{Bmal1}\)-silenced Hepa1-6 cells. Cell survival ranged from 99.6 ± 0.2% after FY26 exposure onset at T22 down to 99.0 ± 0.2% at T04 (ANOVA, \(p=0.194\), Cosinor Analysis: \(p=0.313\)) (Figure VII-15).
Figure VII-15: Time dependent cell toxicity of 1 µM FY26 on Hepa1-6 Per2-luc shBmal1 cells.

The graphs show the cell survival following a 24 h exposure of 1 µM FY26 on Hepa1-6 Per2-luc shBmal1 cell according to different drug administration times. The total protein content was measured 72 h using the photo-colorimetric endpoint measurement SRB assay.

The bioluminescence of the vehicle treated cells, declined 5 days after cell seeding and reached “0” at T16 and T22. Exposing the cells at T04, T10, T16 and T22 to 1 µM FY26 led to a depletion of Per2-luc oscillation approximately 24 h after the start of drug exposure. FY26 decreased Per2-luc bioluminescence (mean ± SEM) from 1812 ± 287 photon counts per second to 440 ± 68 photon counts per second following dosing at T22. Dosing at T04, reduced bioluminescence from 1030 ± 48 photon counts per second to 362 ± 62 photon counts per second (Figure VII-16). Comparing the bioluminescence pattern with the ones observed for Hepa1-6 Per2-luc cells, no increase of bioluminescence was observed 3 to 5 days after the start of drug administration. This further supports previous data that the shBmal1-silenced cell clone is characterised by a higher sensitivity towards FY26 than Hepa1-6 Per2-luc.
Figure VII-16: Time dependent effect of FY26 on the Hepa1-6 Per2-luc shBmal1 bioluminescence.

The AUC$_{Diff}$ was calculated as described for the Hepa1-6 Per2-luc. The bioluminescence curves at T04 and T10 did not cross each other. In order to be consistent with the AUC$_{Diff}$ at T16 and T22, the AUC$_{Diff}$ at T04 and T10 were calculated from 3.4/3.8 days to 5.8 days.

The maximum AUC$_{Diff}$ occurred at T04 was with 4507 ± 173 photon counts per second ∙ h (mean ± SEM) 3 fold higher as the minimum AUC$_{Diff}$ at T22 (1517 ± 237 photon counts per second ∙ h) (Figure VII-17). ANOVA was statistically significant with $p<0.0001$ between the AUC$_{Diff}$ at T04 vs. T10 (***), T16 (***), T10 vs. T16 (**), and between T10 vs. T22 (**). Cosinor Analysis showed a significant 24-h rhythm ($p<0.0001$) with a mesor (mean ± SEM) at 3037 ± 138 photon counts per second ∙ h with an amplitude of 1783 ± 210 photon counts per second ∙ h and an acrophase at $T \pm 0.7 \text{ d}$.
Figure VII-17: Time dependent changes of the AUC\textsubscript{Diff} according to the time of FY26 administration (n=3).

Thus, the maximum Per2-luc bioluminescence inhibition in Hepa1-6 Per2-luc shBmal1 cells was observed at T04 whereas the maximum inhibition in Hepa1-6 Per2-luc cells occurred 6 h earlier at T22.

4.3 Time dependent effect of 1 µM and 2.5 µM FY26 on the Per2-luc reporter construct in HCT116 Per2-luc

The effect of 2.5 µM FY26 and 1 µM FY26 on the Per2-luc bioluminescence in HCT116 Per2-luc cells was assessed as described for the previous experiments. Time dependent changes of the cell survival was observed after the exposure to 0.8 µM FY26 (data taken from the IC\textsubscript{50} chronotoxicity study) with the highest cell survival at T22 with 99.4 ± 0.2% and the lowest at T04 with 97.2 ± 0.3% [ANOVA with p<0.0001 and TMCT: T04 vs. T22 (****), T22 vs. T10 and T16 (****)]. Cosinor Analysis for a 24-h rhythm (p<0.0001) detected a mesor with 98.8 ± 2.5%, amplitude with 11.5 ± 3.7% and an acrophase at T19.9 ± 1.2 h.
Figure VII-18: Time dependent cell survival of HCT116 Per2-luc cells after the exposure of 0.8 µM FY26 at T04, T10, T16 and T22.

Next, the effect of 1 µM or 2.5 µM FY26 at T04, T10, T16 and T22 was assessed on the bioluminescence of HCT116 Per2-luc cells (Figure VII-19). Vehicle exposure did not affect bioluminescence as an increase was observed until day 6 to 7. After day 6 to 7 bioluminescence started to decline except at T16. The administration of 2.5 µM and 1 µM FY26 led to a sharp decline of bioluminescence without any subsequent recovery at all circadian time points except at T10. Here, the bioluminescence of HCT116 Per2-luc cells exposed to 1 µM FY26 recovered. A dose response was observed with a higher inhibition of bioluminescence after the administration of 2.5 µM FY26 as compared to 1 µM FY26.
In order to analysis the time dependent differences, the difference between the median AUC_{vehicle} and the AUC of each bioluminescence curve after the administration of 1 µM and 2.5 µM FY26 were calculated from the start of FY26 exposure until the end of bioluminescence recording (Figure VII-20).

The maximum and minimum of the AUC_{Diff1µM} occurred at T16 with 2732 ± 89 photon counts per second · h and at T04 to T10 with 1843 ± 139 and 1582 ± 603 photon counts per second · h (ANOVA with $p=0.137$, Cosinor Analysis 24-h rhythm with $p=0.192$).

The maximum for the AUC_{Diff2.5µM} occurred at T16 with 2838 ± 44 photon counts per second · h. However, different form the minimum for the AUC_{Diff1µM}, the minimum of the AUC_{Diff2.5µM} was shifted for 6 h to T22 and T04 with 2219 ± 46 and 2227 ± 11 photon counts per second · h. ANOVA was statistically significant ($p<0.0001$) with differences between the time points T04 vs. T10 (***) and T16 (****), between T10 vs. T16 (*) and T22 (***) and between T16 vs. T22 (***)-0.0001). A statistically significant 24-h rhythm ($p<0.0001$) was detected with a mesor of 2475 ±
Results

26 photon counts per second \cdot h, an amplitude of 364 ± 37 photon counts per second \cdot h and an acrophase at T13.8 ± 0.4 h.

Figure VII-20: Results of the difference between the median AUC of the vehicle and each AUC after the administration of 1 µM FY26 and 2.5 µM FY26.

Panel A depicts the results of the AUC_{Diff1µM} and panel B shows the results of the AUC_{Diff2.5µM}.

To sum up, a dose dependent difference between of 1 µM FY26 and 2.5 µM FY26 on the bioluminescence curve was measured. A significant difference was obtained for the AUC_{Diff2.5µM} with the maximum inhibition of bioluminescence at T16. This was consistent with the results of the IC_{50} chronotoxicity as T16 was identified as one of the time point of highest FY26 toxicity together with T04.
5 Postamble

The pharmacokinetics study demonstrated that maximum cellular uptake of FY26 resulted from FY26 exposure starting at T02. Minimum uptake was found for FY26 exposures starting at T10 or T14. Viable cells were also counted in the same dishes as those used for pharmacokinetics determinations. According to the AUC$_{0h-72h}$ and the Slope$_{0h-72h}$, the highest cell viability was observed at T10 and the least at T18. This was further confirmed by the doubling rate as the doubling time was prolonged at T18.

Following the cell viability studies, the effect of FY26 on the cell cycle showed an accumulation of cells in S-phase which was dose dependent. The number of Hepa1-6 Per2-luc cells in G2/M-phase decreased whilst the number of cells in Go/G1-phase increased. This progression was not observed in shBmal1-silenced Hepa1-6 Per2-luc cells as both the number of cells in Go/G1- and in G2/M-phase decreased. This suggests that cells undergo apoptosis rather than to enter G0/G1-phase. The determination of a higher number of apoptotic Hepa1-6 Per2-luc shBmal1 cells supported this observation. Besides apoptosis, the induction of autophagy pointed towards a dose dependent increase of LC3B levels with higher levels detected immediately instead of 72 h after the end of the FY26 incubation time. Determining the time dependent cell cycle phase distribution at T04 and T16 for both cell lines, showed that for Hepa1-6 Per2-luc a higher number of cells accumulated in G2/M-phase at T04. At T16 a higher cell distribution in G0/G1-phase was observed. No such effect was observed for the Bmal1-silenced clone. The highest inhibition of Per2-luc bioluminescence recordings occurred at T22 for Hepa1-6 Per2-luc and at T16 for HCT116 Per2-luc. The maximum Per2-luc inhibition for the shBmal1-silenced cell line was determined at T04.
VIII Results – The effect of FY26 on the circadian clock and the CTS
1 Preamble

The last experiment of the previous chapter shows the time dependent inhibition of bioluminescence after the FY26 administration at different time point. So far, it is unclear whether the inhibition of bioluminescence is caused by an inhibition of cell proliferation or by the interaction of FY26 with the Per2 promoter of the Per2-luc reporter construct and thus with the circadian clock.

Thus, the interaction of FY26 with the circadian clock on a transcriptional translational and on the CTS are presented in this chapter.
2 Effect of FY26 on *Per2-luc* oscillation

The administration of FY26 on Hepa1-6 *Per2-luc* cells at different time points did not disturb *Per2-luc* oscillations (Figure VIII-1) although an initial increase of the amplitude was observed. Nevertheless, Hepa1-6 *Per2-luc* oscillations were detected with a period (mean ± SEM) of 23.9 ± 0.1 h after vehicle exposure and with a period of 25.3 ± 0.8 h after the exposure of 1 μM FY26.

![Figure VIII-1: Detrended bioluminescence of Hepa1-6 Per2-luc after the exposure of media, vehicle, 0.2 μM and 1 μM FY26 at T04, T10, T16 and T22.](image)

The administration of the vehicle to the Hepa1-6 *Per2-luc shBmal1* cells did not alter *Per2-luc* bioluminescence oscillations. However, the introduction of 1 μM FY26 dampened the amplitude of *Per2-luc* bioluminescence at all 4 time points (Figure VIII-2). The mean ± SEM period after vehicle administration was 23.4 ± 0.9 h and after FY26 administration 22.5 ± 1.6 h thus confirming the dampened amplitude of *Per2-luc* oscillations after the exposure of 1 μM FY26.
Results

Figure VIII-2: Detrended bioluminescence of Hepa1-6 Per2-luc shBmal1 bioluminescence after the exposure of vehicle and 1 µM FY26 at T04, T10, T16 and T22.

The bioluminescence of the Per2-luc reporter in HCT116 cells was not altered after vehicle administration. The mean ± SEM period was calculated as 24.4 ± 0.6 h under the vehicle exposure. However, after the administration of 1 and 2.5 µM FY26, amplitudes were dampened except at T16 and at T10 for 1 µM FY26. The mean ± SEM periods were determined with 21.9 ± 0.9 h for 1 µM FY26 and with 23.2 ± 0.4 h for 2.5 µM FY26. This suggests that FY26 effected cell proliferation rather than the Per2-luc reporter.

To summarise, the administration of FY26 did not alter the Hepa1-6 Per2-luc oscillation but did effect the amplitudes of the shBmal1-silenced Hepa1-6 clone and of HCT116 Per2-luc. Whether this effect was caused by an interaction of FY26 with the circadian clock will be investigated in the following chapter using Hepa1-6 Per2-luc cells.
Results

**Figure VIII-3**: Detrended bioluminescence curves of HCT116 Per2-luc after the exposure of vehicle, 1 µM and 2.5 µM FY26 at T04, T10, T16 and T22.

### 3 Effect of FY26 on clock gene expression

Circadian patterns in mRNA expression of clock genes *Per2*, *Bmal1* and *Rev-erba* were determined in temperature synchronised Hepa1-6 *Per2-luc* cells with or without exposure to 1 µM FY26. The drug was added at T04 and at T22 (Figure VIII-4).

No difference between the *Per2* gene expression after the vehicle administration at T04 and at T22 was observed. The relative gene expression (RGE) was identified with 0.38 ± 0.09 RGE (mean ± SEM) for T04 and with 0.29 ± 0.03 RGE for T22. The acrophase of the *Per2* gene expression at T04 and at T22 was determined at T 15.6 h (Cosinor, \( p<0.001 \)) and at T 13.4 h (Cosinor, \( p=0.125 \)). No changes were observed following the FY26 administration at T04 and at T22. The highest *Per2* expression following FY26 administration occurred at T16 with 0.28 ± 0.13 RGE for T04 and with 0.18 ± 0.04 RGE for T22. The acrophase of the *Per2* expression of with FY26
incubated cells at T04 and T22 was calculated with T15.7 h (Cosinor, $p=0.406$) and with T 14.4 h (Cosinor, $p=0.505$) (Figure VIII-4 and Table VIII-1).

Following the vehicle administration at T04 and at T22, $Bmal1$ peak occurred at T20 with 1.27 ± 0.49 RGE (mean ± SEM) and T22 with 1.27 ± 0.49 RGE. The acrophase was calculated with T16.2 h (Cosinor, $p=0.614$) for T04 and with T17.8 h (Cosinor, $p=0.757$). The FY26 exposure did not alter the $Bmal1$ gene expression. The highest $Bmal1$ expression occurred at T20 with 1.29 ± 0.39 RGE following dosing at T04 and at T22 with 1.8 ± 1.5 RGE after FY26 exposure at T22. The acrophases were calculated with T21.9 (Cosinor, $p=0.022$) for T04 and with T18.7 (Cosinor, $p=0.608$) for T22 (Figure VIII-4 and Table VIII-1).

The highest expression of the clock gene Rev-erba under the vehicle exposure was observed at T22 with 0.52 ± 0.18 RGE for T04 and at T16 with 1.55 ± 0.29 RGE for cells exposed to the vehicle at T22. Cosinor Analysis was applied and identified with an acrophase for Rev-erba expression at T16.9 h ($p=0.016$) and at T14.9 h ($p=0.231$) for the vehicle administration at T04 and at T22. However, dosing cells at T04 with FY26 altered the maximum of Rev-erba expression from T22 to T12 with 0.82 ± 0.52 RGE. No change of Rev-erba expression was observed when FY26 was administered at T22. The relative gene expression was thus highest at T16 with 2.12 ± 0.31 RGE. Nevertheless, Cosinor Analysis identified the acrophase at T15.2 h for T04 ($p=0.339$) and at T15.8 for T22 ($p=0.088$) (Figure VIII-4 and Table VIII-1).

To conclude, exposing 1 µM FY26 to Hepa1-6 Per2-luc cells did not affect the expression of the clock genes Per2, Bmal1 and Rev-erba 20 h to 24 h after FY26 administration at T04 and T22 as their pattern was similar to the gene expression pattern after vehicle administration. This suggest that the decline of photon counts per second was not related to the suppression of gene expression by FY26 but rather to the cytostatic effect of FY26 on Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 and HCT116 Per2-luc.
Figure VIII-4: Effect of 1 µM FY26 on the gene expression of the clock genes Per2, Bmal1 and Rev-erba whilst synchronised with temperature cycle D at T04 (left panel) and at T22 (right panel).
Table VIII-1: Cosinor Analysis of the clock gene expression after the exposure of the vehicle and FY26 at T04 and T22.

The statistically significant results are highlighted in bold.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Treatment</th>
<th>Period</th>
<th>Mesor in RGE</th>
<th>Amplitude in RGE</th>
<th>Acrophase in time (h)</th>
<th>p-value</th>
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<td><strong>Per2</strong></td>
<td>Vehicle</td>
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<td></td>
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<tr>
<td>FY26</td>
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<td>4.1</td>
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<tr>
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<td>16.2</td>
<td>0.614</td>
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<tr>
<td></td>
<td>12-h</td>
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<td></td>
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<tr>
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<td>0.5 ± 0.1</td>
<td>21.9</td>
<td>0.022</td>
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<td>3.6</td>
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<td>12.2</td>
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<td>0.79 ± 0.16</td>
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<td>0.2</td>
<td>18.7</td>
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<td>8.2</td>
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</tr>
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<td><strong>Rev-erba</strong></td>
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<td>0.2</td>
<td>14.9</td>
<td>0.231</td>
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<td></td>
<td>12-h</td>
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<td></td>
<td>0.2 ± 0.1</td>
<td>3.4 ± 0.6</td>
<td>0.021</td>
</tr>
<tr>
<td>FY26</td>
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<td>0.6 ± 0.1</td>
<td></td>
<td>0.4</td>
<td>15.8</td>
<td>0.088</td>
</tr>
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<td></td>
<td></td>
<td>0.05</td>
<td>3.0</td>
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</tbody>
</table>
4 FY26 affects circadian timing system biomarkers in mice

4.1 FY26 disrupts mouse core body temperature if dosed at ZT18

Six C57Bl/6 mice were subjected to antiphasic LD12:12 synchronisation immediately after the implantation of the temperature sensor into the peritoneal cavity. Core body temperature was automatically recorded at 5-min intervals throughout the whole experiment. Up to 8 days were needed for the mice in the antiphasic LD schedule to adjust the core body temperature cycle to the new synchroniser. Thus, all the mice were well synchronised by their respective LD12:12 schedules after 3 weeks (data not shown).

The mice in both groups were intraperitoneally injected at ZT06 or at ZT18 with vehicle, then with 50 mg/kg FY26 one week later. FY26 treatment induced up to 8% body weight loss 24 h after dosing in individual mice (Figure VIII-5).

Figure VIII-5: Time dependent body weight loss after a single i.p. injection of the vehicle and FY26.

Panel A: body weight loss after a single i.p. injection of the vehicle at ZT06 and ZT18. Panel B: Body weight loss after a single i.p. injection of 50 mg/kg FY26 at ZT06 and ZT18 (n=3 per circadian time point).

Maximum body weight loss (mean ± SEM) was 3.6 ± 1.2% following drug dosing at ZT06, as compared to 5.3 ± 1.7% at ZT18 (two-sample t-test for equal mean with
Results

The effects of vehicle and FY26 on core body temperature were determined in the same mice as a function of dosing time (Figure VIII-6). Vehicle injections did not affect core body temperature circadian patterns. In contrast, an immediate and sharp decline was observed in the mice injected at ZT18, with temperature acutely dropping by 3.7°C (from 37.5 ± 0.2°C to 33.8 ± 0.3°C) at 6 h post injection. In mice treated at ZT06, temperature decreased only by 1°C (from 35.7 ± 0.7°C to 34.7 ± 0.5°C), and such nadir was reached 2 h after injection. The differences between the extents of temperature decline according to FY26 timing were statistically significant ($t$-test with $p=0.006$).

Figure VIII-6: Effect of a single intraperitoneal injection of the vehicle and 50 mg/kg FY26 on core body temperature in mice.

The panel depicts the average of the hourly average core body temperature of three mice at ZT06, ZT18 and of 6 mice for the vehicle.

To further analyse the effects of vehicle and FY26 on core body temperature patterns, hourly averages were computed from the 5-min temperature records. The period was estimated for three days before and after FY26 administration using spectrum resampling algorithm as shown in the appendices Figure XI-18 and Figure XI-19.
Estimated periods in individual mice were averaged over the 24-h span after vehicle or FY26 administration (Figure VIII-7). The endogenous period of the temperature rhythm remained stable at 22.8 ± 0.3 h or 22.9 ± 0.2 h in the mice receiving vehicle at ZT06 or ZT18 respectively. FY26 administration significantly lengthened the circadian period by 0.7 ± 0.5 h following dosing at ZT06 (23.5 ± 0.4 h), but it shortened it by 2.4 ± 0.4 h in the mice treated at ZT18 (20.6 ± 0.6 h). ANOVA revealed a significant difference of the periods according to timing (p=0.01) but not between the difference treatments (p=0.06). However, the interaction between time and treatment was significant with 0.004. Tukey’s multiple comparison showed a significant difference between the periods of the vehicle treated mice at ZT18 vs. the FY26 treated at ZT18 (*) and between the period of the FY26 treated mice at ZT06 and ZT18 (**).

*Figure VIII-7: Period of mice core body temperature 24 h after the injection of the vehicle and 50 mg/kg FY26 at ZT06 and ZT18.*

The results thus demonstrated that FY26 treatment did not affect the core body temperature rhythm when treated at ZT06. However, a drop of core body temperature by 3.7°C and a shortening of the temperature period by 2.4 h was encountered following dosing at ZT18. The results suggested a pronounced time dependency of FY26 transient effects on the CTS, with largest rhythm alterations corresponding to FY26 dosing at ZT18, i.e. the timing yielding the worse systemic toxicity.
4.2 The effect of FY26 on the PER2::LUC liver expression

Nine C57Bl/6 mice were first synchronised with LD12:12, then had a temperature sensor implanted into the peritoneal cavity and the posterior back skin shaved. Each mouse was then singly housed in the RT-Bio with luciferin-containing bottles and kept in DD for 7 to 13 days. After 3 to 5 days of baseline recording, three mice received a single i.p. vehicle injection at CT06 or CT18, while three mice were injected with 50 mg/kg FY26 at CT06. Three other mice received the same treatment at CT18. As mice were kept under constant darkness, the period advanced every 24 h by 1.5 h. Injections being times at CT06 and CT18 referred thus to CT22.5-CT01.5 and to CT10.5-CT13.5 instead.

No body weight loss was observed after vehicle administration. In contrast, body weight loss (mean ± SEM) was largest 24 h after FY26 treatment irrespective of drug timing, being 6.6 ± 1.6% and 5.8 ± 4.3% for FY26 dosing at CT06 or CT18 respectively (p from ANOVA = 0.279) (Figure VIII-8).

![Figure VIII-8: Body weight changes after FY26 dosing. Vehicle or 50 mg/kg FY26 were injected in one single i.p. injection at CT06 and CT18 in 9 individual PER2::LUC mice.](image)

The effects of vehicle or FY26 on PER2::LUC expression time series were analysed using Spectrum Analysis for the data sets from each of the 9 mice. Individual profiles of hourly averaged normalised photon counts per min including the period, amplitude and acrophase are shown in Appendix Figure XI-20. Figure VIII-9 depicts the average PER2::LUC bioluminescence for the 2 days preceding and the 3 days following vehicle or FY26 dosing at putative CT06 or CT18 respectively, under the assumption
of the persistence of 24-hour periodicities in DD. The vehicle injection at CT06 and at CT18 did not affect PER2::LUC bioluminescence. The administration of 50 mg/kg FY26 led to the ablation of PER2::LUC oscillations at CT06 and CT18 which recovered after 1.5 days for CT06 and 2 days for CT18. Interestingly, no disruption of PER2::LUC oscillations following FY26 dosing at CT06 and CT18 were observed for mouse 1 and 3.

Figure VIII-9: Effect of a single injection of the vehicle and 50 mg/kg FY26 at CT06 and CT18 on the PER2::LUC expression in PER2::LUC mice.

The graph depicts the mean of the hourly average of the normalised photon counts per min that reflect PER2::LUC expression in 3 mice receiving i.p. vehicle at CT06 (panel A), vehicle at CT18 (panel B) or 50 mg/kg FY26 at intended CT06 (panel C) or at intended CT18 (panel D).

The mean ± SEM period of PER2::LUC expression remained stable at 22.5 ± 0.3 h after vehicle administration. Following the administration of FY26 at intended CT06 and CT18, PER2::LUC oscillating bioluminescence pattern were abolished.
Differences in bioluminescence patterns were further explored through computing AUC’s over the 24 h timespan after injection. The mean ± SEM AUC’s, in hourly averaged photon counts/min · h, were 10.0 ± 1.5 and 22.1 ± 1.2 following FY26 injection at CT06 or CT18 respectively (p-value from t-test = 0.004) (Figure VIII-10). Thus, the injection of 50 mg/kg FY26 led to a larger decrease of PER2::LUC bioluminescence at intended CT06 as at CT18.

Figure VIII-10: Period and AUC of the PER2::LUC bioluminescence pattern after vehicle and FY26 treatment at intended CT06 and CT18.

Panel (A) shows the change of the period of the PER2::LUC bioluminescence pattern and panel (B) shows the AUC of the PER2::LUC bioluminescence pattern after FY26 administration at intended CT06 and CT18.

Core body temperature was measured simultaneously with PER2::LUC bioluminescence expression. A technical failure happened for the three first mice kept under DD. Thus Figure VIII-11 displays core body temperature rhythms for two mice per condition. Vehicle administration at intended CT06 and CT18 did not alter core body temperature rhythms as compared to baseline. In contrast, FY26 profoundly altered the temperature rhythm at least for 48 h, with a drop by 0.8°C within 12 h and 5 h after the injection at intended CT06 or CT18 respectively. Core body temperature rhythms recovered 2 days after treatment (Appendix Figure XI-21) (Figure VIII-11).
Results

Figure VIII-11: Time dependent changes of body temperature circadian rhythm of PER2::LUC mice treated with 50 mg/kg FY26 or vehicle.

Panel A shows the effect of the vehicle administration at CT06 and CT18 on core body temperature and panel B shows the FY26 effect on core body temperature following dosing at CT06 and panel C at CT18.

The period (mean ± SEM) of the temperature rhythm after vehicle was 23.4 ± 0.3 h. FY26 administration led to the ablation of core body temperature rhythm, whose dominant mean (± SEM) periods were computed at 29.5 ± 3.2 h and 33.9 ± 4.1 h for treatment at intended CT06 and CT18 respectively (Figure VIII-12).

To summarise, FY26 transiently suppressed the circadian rhythm in core body temperature following dosing at ZT18, but not at ZT06 in mice synchronised with LD12:12. The disruption of this circadian biomarker was observed following FY26 dosing at intended at CT06 and CT18 in mice kept in DD for 3 to 5 days before treatment application. Since the average endogenous period of PER2::LUC expression (which variable) in the DD control mice ranged from 22.0 h to 23.0 h, rather than remaining close to 24 hours, intended CT06 thus corresponded to CT22.5 to CT01.5, while intended CT18 was effectively located between CT10.5 and CT13.5. Therefore, the largest reduction in the AUC of the hourly averaged photon counts per min resulted from FY26 dosing at endogenous CT22.5 to CT01.5. This was consistent with the in
vivo chronotoxicity study, where ZT22 was identified as close to the time point of worst FY26 tolerability.

*Figure VIII-12: Period of the mice core body temperature after the administration of vehicle and 50 mg/kg FY26 at intended CT06 and CT18.*
5 Postamble

The *in vitro* studies demonstrated that FY26 did not affect the clock gene expression suggesting that the time dependent decrease of *Per2-luc* bioluminescence is related to FY26 cytotoxic.

Further *in vivo* studies showed that FY26 could alter biomarkers of the circadian timing system at whole organism and molecular levels. The third experiment revealed that a single injection of FY26 (50 mg/kg) led to a sharp decrease in core body temperature thus a transient circadian disruption following treatment at ZT18 but not at ZT06.

The forth experiment showed that the same dose of FY26 disrupted the circadian rhythms in both PER2::LUC bioluminescence for 2 out of 3 mice injected at CT06 and CT18. However, the impact of the disruption was less following dosing at CT18 as the AUC of the PER2::LUC bioluminescence was still higher as compared to CT06. Core body temperature rhythm of mice kept in DD further showed disrupted rhythms which recovered 2 days post treatment.

Taken together, the studies not only showed that FY26 could respect or disrupt biomarkers of the circadian timing system according to its dosing time in light-dark synchronised mice, but also revealed that entrainment by a 12:12 h photoperiod plays a crucial role in the effect of FY26 on the CTS. Comparing the effect of FY26 on core body temperature, demonstrated that FY26 did not affect core body temperature rhythms of mice entrained to LD 12:12 cycles whereas this effect was diminished in mice kept under 12:12 DD.
IX Discussion
Towards the translation of *in vitro* into *in vivo* FY26 chronotoxicity through temperature cycles resetting of circadian clocks

Here, I first discuss the relevance of temperature synchronisation for assessing the *in vitro* chronotoxicity, using the anticancer drug candidate FY26 as an example (chapter IV).

For the *in vitro* model, a murine hepatocarcinoma cell line with and without silenced *Bmal1* and a human colon cancer cell line were chosen. Temperature cycle starting with a 5°C temperature change after the first 12 h followed by alternating temperature changes of 1°C between 36 and 37°C every 12 h, was designed to simulate mouse core body temperature, thus imposing a quasi-physiological temperature rhythm on a cell population (Brown *et al.*, 2002). This temperature cycle “D” synchronised Hepa1-6 *Per2-luc* and HCT116 *Per2-luc* to a robust circadian period (~ 24 h), with phase and amplitude remaining stable for 6 to 7 days. It outperformed other temperature schedules that were assessed regarding period, acrophase and amplitude stabilities over 6 to 7 days. The synchronisation properties of temperature cycle D was further tested by exposing Hepa1-6 *Per2-luc* to T-cycles with 20-h or 26-h periods in order to assess the synchronisation properties. A synchronisation, leading to a stable *Per2-luc* bioluminescence period of ~ 20 h, was achieved with temperature alternating from 37 to 36°C and the reverse every 10 h. Such adequate synchronisation likely reflected the fact that the imposed 20-h period of the synchroniser matched the endogenous circadian period of Hepa1-6 *Per2-luc* cells that was determined in cell cultures released at a constant temperature of 37°C, after an initial 24-h temperature synchronisation flash. Exposing Hepa1-6 *Per2-luc shBmal1* cells to temperature cycle D, resulted in “pseudo-rhythmic” oscillations with a triangle shaped *Per2-luc* bioluminescence pattern. Once Hepa1-6 *Per2-luc shBmal1* cells were released at a constant temperature of 37°C, no sustained oscillations were observed as compared to the clock-proficient cell clone Hepa1-6 *Per2-luc*. This might indicate that the bioluminescence oscillations of Hepa1-6 *Per2-luc shBmal1* under temperature synchronisation could relate to some residual clock function, as 36% *Bmal1* mRNA expression still remained in the *shBmal1* clone. Besides synchronising a defective
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clock in Hepa1-6 Per2-luc shBmal1, temperature cycles might have driven other cellular proteins resulting in triangle shaped Per2-luc peaks. One possible protein could be the heat shock protein 1 (HSP1) as heat shock proteins are upregulated in response to elevated temperature and in response to oxidative stress especially in cancer cells (Oksala et al., 2014; Santos et al., 2017; Werner et al., 2007). The activation of the heat shock factor 1 (HSF1) is regulated by the HSF cycle (Morimoto & development, 1998). The HSF1 has been linked to the circadian clock as it induced the expression of heat shock proteins at the beginning of the active phase in mice. Thus, the expression of HSP followed a circadian pattern (Reinke et al., 2008; Tamaru et al., 2011). The acrophase of the Per2-luc expression in the Bmali silenced cell line Hepa1-6 Per2-luc shBmal1 occurred during the end of the 36°C temperature cycle before the temperature raised to 37°C. This timing corresponded to the beginning of the active phase in mice, when the endogenous temperature rises as a result of its circadian regulation by the SCN. This hypothesis was further supported by the induction of the heat shock protein response following a 1.5°C temperature rise from 37 to 38.5°C. Such response was inhibited by the HSP1 antagonist KNK437 (Buhr et al., 2010). Interestingly, the expression of Hsp reportedly remain rhythmic after the disruption of the circadian clock. In LAP-tTA|TRE-Rev-erva double transgenic mice the molecular clock was disrupted by inhibiting Bmali through the overexpression of REV-ERBα. This resulted in a circadian accumulation of Per2 and Hsp mRNA indicating these rhythms could arise independently from the genetic clock (Kornmann et al., 2007). These findings might well explain the triangle shaped Per2-luc patterns in Hepa1-6 Per2-luc shBmal1 cells, as those cells were exposed to rhythmic external temperature cues. Other proteins which responded to temperature changes are the cold induced proteins RNA-binding proteins CIRBP and RBM3. It has been reported that CIRBP was upregulated in response to cold temperatures (Nishiyama et al., 1997). In mice kept in constant darkness following the entrainment to LD 12:12 cycles, Hsps were upregulated at CT18, during the activity phase, and Rbps were highly expressed in response to lower temperature at CT06 (Liu et al., 2013). The depletion of Cirbp and Rbm3 in mouse fibroblast reduced the amplitude of the clock genes Per1, Dbp, Nr1d1 and Nr1d2, following temperature synchronisation. Further, the binding of genes to PAS were regulated by CIRBP and RBM3 in mouse liver. This resulted in strong circadian oscillations indicating that circadian gene expressions is controlled
by Cribp and Rbm3 under temperature synchronisation (Liu et al., 2013). Further it was reported that the expression of Cribp was regulated by temperature dependent splicing, which was highest at low temperatures (Gotic et al., 2016). Taken together, these data suggest that synchronisation through temperature cycles might be independent from the circadian clock, thus accounting for the “pseudo-rhythmicity” observed in temperature synchronised Hepa1-6 Per2-luc shBmal1.

I have further demonstrated that the expression of core clock genes Per2, Bmal1 and Rev-erba in clock-proficient Hepa1-6 Per2-luc cells were rhythmic. The sampling of the cell pellet for RNA extraction was done 2.5 days after the start of temperature synchronisation. This time window was chosen as stable period lengths, acrophases and amplitudes were achieved 2.5 days after the start of synchronisation (data not shown). The Bmal1 peak was determined at T0, when the temperature increased from 36°C to 37°C. The Per2 peak occurred at T12 following the programmed temperature decrease from 37 to 36°C, whereas the acrophase of the Rev-erba expression occurred at T16, i.e. 4 h after the Per2 peak and 8 h before the Bmal1 peak. The bioluminescence curves of the Per2-luc and Bmal1-luc reporters into the HCT116 cells, showed antiphase oscillation patterns, whilst being subjected to the same temperature cycle D. Both oscillations persisted and remained in antiphase, once the cells were released at a constant temperature of 37°C, as it was reported for the Hepa1-6 Per2-luc cells. Thus, synchronising both cell lines with a 24-h temperature cycle has revealed autonomous and self-sustained circadian clock genes expression patterns, using bioluminescence reporter technology in line with other reports (Nagoshi et al., 2004).

The implication of circadian clock coordination for cell cycling was then examined in synchronised Hepa1-6 cell populations, since a 1:1 coupling between both oscillators was demonstrated in fibroblasts (Bieler et al., 2014; Feillet et al., 2014). Alternating 12:12 h temperature cycles synchronised S-phase, but neither G0/G1 nor G2/M phase cells, to a 24-h period in Hepa1-6 Per2-luc. Hepa1-6 Per2-luc shBmal1, however, showed a 24-h rhythm in G0/G1-phase and a 12-h rhythm for G0/G1- and S-phase. This suggested that the cell cycle was at least partly uncoupled from the circadian clock in this model. Several studies have investigated the relationship between the circadian clock and the cell cycle (Bieler et al., 2014; Feillet et al., 2014). Interestingly, such
coupling was lost in some cancerous cell lines as described in the doctoral thesis of Elham (Aida) Farshadi. She demonstrated that the cell cycle in mouse breast carcinoma cells MBC399 was uncoupled from the circadian clock which was free-running (Farshadi, 2018). In my studies, the cell cycle phases distribution were determined under 24-h temperature synchronisation of the circadian clock as indicated with stable ~24 h oscillations of the Per2-luc reporter. Yet, the data support a partial uncoupling of the cell cycle in this model that would need further exploration using simultaneous clock and cell cycle phase imaging reporters.

Temperature schedule D was then used to assess any time dependent toxicity of FY26 in synchronised cell populations (chapter III.2). Toward this goal, the in vitro chronotoxicity of FY26 was determined in Hepa1-6 Per2-luc, Hepa1-6 Per2-luc shBmal1 and HCT116 Per2-luc. The IC50 of FY26 in Hepa1-6 Per2-luc cells was significantly highest following drug exposure at T22, suggesting this to be the time point of least drug toxicity. Thus at T22, a higher FY26 concentration was required to achieve a cell survival of 50% as compared to other times of FY26 addition. Whereas the IC50 of Hepa1-6 Per2-luc increased following drug addition from T04 to T22, the IC50 of FY26 in Hepa1-6 Per2-luc shBmal1 depicted a bimodal pattern identifying T10 and T22 as time points with the least FY26 toxicity. In schedule D, T10 and T22 are situated 2 h before the temperature decreases from 37 to 36°C and temperature increases from 36 to 37°C, respectively. The IC50 of FY26 in HCT116 Per2-luc cells also exhibited a bimodal pattern although being clock-proficient. As observed in Hepa1-6 Per2-luc shBmal1, the highest IC50 was determined at T22 and the second highest value at T10. This suggested that the bimodal FY26 toxicity pattern might not only be related to a genetically disrupted circadian clock, but also to tissue specific entrainment properties and/or to the clock regulation of FY26 detoxification mechanisms.

The relevance of in vitro chronotoxicity data to predict for in vivo chronotoxicity patterns was investigated in chapter III.4, using two osmium complexes, FY26 and FY25. FY26 and its active metabolite FY25 (Peter Sadler, personal communication) were administered intraperitoneally at one of six equally spaced dosing times over the 24 hours to male C57Bl/6 mice synchronised with LD12:12, where ZT0 and ZT12 correspond to light onset and offset respectively. FY26, but not FY25, exhibited a
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statistically significant circadian toxicity profile, based on body weight changes following drug administration. Maximum body weight loss was highest following FY26 treatment during the second half of the active phase (ZT18-ZT22) and lowest in mice dosed at ZT02, near the beginning of the inactive phase. The time points of highest and lowest toxicity were further confirmed by the assessment of the AUC of body weight loss over the 7 days following drug administration \((\text{AUC}_{0d-7d})\) and by the slope of body weight recovery to pretreatment body weight. The highest \(\text{AUC}_{0d-7d}\) occurred at ZT02 and the lowest at ZT22. Mouse body weights recovered fastest following FY26 injection at ZT06 and slowest after receiving it at ZT18. Cosinor Analyses revealed significant circadian rhythms thus identified the acrophases of least body weight loss, and \(\text{AUC}_{0d-7d}\) and that of fastest recovery slope (best tolerability). The acrophases of these three parameters occurred at ZT6.7, ZT6.1 and ZT6.2 respectively. This indicated that FY26 was best tolerated if administered at ZT6, i.e. near the middle of the inactive rest phase of the rest-activity circadian cycle.

So far no chronotoxic profile had been reported for any osmium compound. However, the chronotoxicity patterns of platinum anticancer complexes have been thoroughly researched since the initial fortuitous discovery of cisplatin antitumour activity by Barnett Rosenberg (1969). Cisplatin was best tolerated when given near ZT15 to ZT18 to mice or rats (Boughattas et al., 1990; Hrushesky et al., 1982b). As demonstrated for cisplatin, carboplatin and oxaliplatin were also best tolerated following treatment at ZT15 to ZT16 in mice. Worse renal and hematologic dose limiting toxicities of cisplatin, and worse hematologic and intestinal dose limiting toxicities of carboplatin and oxaliplatin were achieved in rodents treated between ZT6 to ZT10 (Boughattas et al., 1994; Boughattas et al., 1990; Boughattas et al., 1989; Ron et al., 1998).

However, following up to six i.p. injections of FY26 (subacute toxicity) did not result in any histopathological changes, although mortality was 12 to 17%. This questioned the mechanisms involved in the toxicity of this metallo-compound. One of the ligands of FY26 was iodide which was replaced by chloride after its cellular uptake, being activated to FY25. Based on an oral communication with Professor Peter Sadler, 1 mg FY26 releases 150 µg iodide. As mice were injected with 50 mg/kg FY26 corresponding to 1.5 mg FY26 for a 30 g mouse, 225 µg iodide were thus co-administered. The LD50 doses of i.p. potassium iodide and iodate were reported to
be 1117 ± 30 and 136 ± 5 mg/kg respectively (Webster et al., 1957). This refers to 33.5 mg iodide and 4.1 mg iodate for a 30 g mouse. Thus, repeated injections of FY26 could result in iodate intoxication. FY26 also increases ROS, thus hydrogen peroxide (H$_2$O$_2$); H$_2$O$_2$ can react with excess iodide to iodate and exceed the iodate LD$_{50}$ (Schmitz, 2001). FY26 lethal toxicity in mice could hence result from acute iodide toxicity, which is known to involve necrosis as well as cytoplasmic fragments desquamation, ER vesiculation and accumulation of lipofuscin in secondary lysosomes in thyroid cells (Many et al., 1992).

Our data revealed that FY26 was best tolerated following dosing in the middle of the rest phase (ZT6), i.e. nearly 12 h apart from the optimal timing of platinum complexes, as well as that of Cd or Hg salts (Cal. J. C., 1984; Miura et al., 2012). FY26 was hence the first metallo-compound whose best tolerability resulted from its delivery during the rest phase of rodents. Thus the results from chronotoxicology studies supported the occurrence of clearly different pharmacologic mechanisms between FY26 and most other metallo-compounds, one of which could be related to acute iodide release.

The consistency of the in vitro and in vivo chronotoxicity patterns of FY26, was further shown using the 24-h rhythm in temperature exposure as a reference biomarker. Best tolerability of FY26 occurred following drug addition at T18 to T22 in temperature synchronised cell cultures, and at ZT06 in LD12:12 synchronised mice. The time points of best FY26 tolerability in vitro and in vivo thus referred to the acrophases of both temperature cycles as depicted in Figure IX-1. Taken together, the results demonstrated that the endogenous temperature rhythms was an adequate circadian biomarker for predicting FY26 treatment in vivo based on the in vitro experiments. Core body temperature in humans is in antiphase to that in mice (Morf & Schibler, 2013). Thus, FY26 would hypothetically be best tolerated in humans when given near the middle of the night, whereas cisplatin and oxaliplatin were best tolerated when administered during the late afternoon, at 4 to 6 pm (Levi et al., 1990).
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Figure IX-1: Schematic representation of the temporal relations between the circadian rhythm in temperature and FY26 chronotolerance in vitro and in vivo. Results from Cosinor Analysis of the endogenous (dotted blue line) temperature cycle and core body temperature rhythm (dotted red line).

The graph presents the 24-h periodic cosine curves best fitting the temperature cycles measured in cell media-containing dishes exposed to exogenous temperature schedule D (blue), and in LD 12:12 synchronised mice (Chapter V). Both curves are aligned in phase.

2 Anticancer properties of FY26 at the time points of highest and lowest tolerability

The balance between antitumor efficacy and tolerability is a critical issue in cancer chemotherapy. FY26 displayed modest efficacy against Hepa1-6, in nude CD1 mice. Dose-response experiments conducted at ZT06 first revealed that doses of 40 or 60 but not 80 mg/kg were in the therapeutic range. Mice dosed with 40 rather than 60 mg/kg FY26 showed a higher tumour growth inhibition. This is striking as the opposite effect was expected. According to the data showing a higher tumour efficacy at 40 mg/kg as at 60 mg/kg FY26, FY26 efficacy could display a bell-shaped tumour efficacy profile. A bell-shaped efficacy profile has so far been reported for the opioid buprenorphine. At lower doses buprenorphine exhibited a high antinociceptive effect which decreased at higher doses. Such effect is caused as buprenorphine has two
distinct pharmacological targets: it is both an agonist of the µ-receptor and a
cot-activator of the opioid-receptor like 1 (ORL-1) receptor (Lizasoain et al., 1991; Lutfy et al., 2003). Such unusual dose response relation for an anticancer agent needs confirmation on a larger sample size. Yet, the buprenorphine example indicates that a bell-shaped FY26 efficacy profile is possible if FY26 has two cellular targets with different susceptibility thresholds. A more frequent dose-response pattern for efficacy is of an exponential type, with a “ceiling” effect for efficacy. For instance, several studies reported that cisplatin did not exhibit differences in efficacy between “low” (40 to 50 mg/m$^2$) and “high” (75 to 100 mg/m$^2$) dose levels in patients with head and neck cancer (Szturz et al., 2017; Wong et al., 2015). A meta-analysis comparing low dose versus high dose chemotherapy reported that anticancer efficacy was not impaired at low doses, which significantly reduced toxicity (Xie et al., 2017). These studies pointed towards a possible ceiling-effect of anticancer agents whose dose reduction might lower side effects but not efficacy.

Next, the time dependent tumour growth inhibition following dosing with 50 mg/kg FY26 at ZT06 and ZT18 was tested. The FY26 dose of 50 mg/kg was chosen based on the previous therapeutic dose selection study that did not detect any statistically significant difference in efficacy between 40 and 60 mg/kg FY26. ZT06 and ZT18 had been identified as the time point of best and worst tolerability respectively. Tumour growth delay was ~40% larger at ZT18 as compared to ZT06, suggesting timing could influence FY26 antitumor efficacy. For the platinum compounds cisplatin and oxaliplatin, tumour growth inhibition was highest following dosing at ZT15 (Granda et al., 2002; Li et al., 2005a; Sothern et al., 1989). Thus, either platinum compounds exhibited highest efficacy following dosing at their best tolerated time points. FY26 on the contrary, seemed to inhibit tumour growth more following its administration at the time point of worst toxicity, under the alternate day schedule that was selected. The anticancer drugs are also best tolerated during the light/ (rest) span in male mice, like FY26. This is the case for both topoisomerase inhibitor irinotecan and the mitotic spindle poison docetaxel for FY26. However, both drugs were also most effective following their administration near their respective circadian times of best tolerability. Interestingly though, docetaxel efficacy was highest following dosing at ZT7 on a
The results from both efficacy studies put FY26 in a special position compared to the above mentioned anticancer agents. As both studies involved a limited number of mice, additional studies testing different FY26 dosing schedules in different tumour-bearing models, with adequate statistical power, are needed to confirm the current results. Further investigations of the mechanisms of action of FY26 are also indispensable as this might explain the differences in time dependent toxicity and efficacy profiles.

3 The circadian clock effects the pharmacokinetics and pharmacodynamics profile of FY26

3.1 Chronopharmacokinetics of FY26

FY26 exhibited a circadian toxicity profile with best tolerability in the middle of the rest phase when core body temperature was lowest. The interplay between the circadian clock and FY26 pharmacokinetics and toxicity is discussed here.

The in vitro chronopharmacokinetics study revealed that maximum intracellular osmium concentrations resulted from FY26 addition at T02 and minimum ones at T14. The area under the curve of the graph depicting the intracellular osmium concentration curve from the time of drug introduction to $t_{max}$ (AUC$_{0-h-t_{max}}$) was highest at T02 and lowest at T10 with a 2-fold difference. Cosinor analysis further confirmed the time points of maximum FY26 uptake with acrophases of the three selected endpoints being located near T02. Thus FY26 uptake displayed a circadian pattern with highest value at T02 and lowest one at T14. These times corresponded to 2 h after temperature increase from 36 to 37°C and decrease from 37 to 36°C respectively. According to the in vitro chronotoxicity data using IC$_{50}$ as an endpoint, T22 had been identified as the time point of lowest toxicity. Thus, there was an 8 h time lag between the times of least FY26 uptake at T14 and that of least toxicity at T22. Interestingly, the highest FY26 uptake at T02 was only 2 h apart from the time point of worst toxicity at T04 suggesting FY26 to depict an asymmetry toxicity pattern. Overall the results indicated that the chronopharmacokinetics and chronopharmacodynamics patterns of FY26 did
not overlap. Such differences between circadian pharmacokinetics and pharmacodynamics profiles had been emphasised for the β-blocker propranolol chronopharmacology in human subjects. Maximum and minimum plasma concentrations were detected following oral propranolol administration at 08:00 and at 02:00 respectively. In contrast, propranolol efficacy, as assessed with heart rate decrease, followed no circadian pattern, thus indicating the dissociation between chronopharmacokinetics and chronopharmacodynamics could indeed be sharp (Langner & Lemmer, 1988).

No transporter mechanism has been identified so far for FY26. However, according to a first systems pharmacology study, FY26 uptake involved energy dependent transport (Doctor Annabelle Ballesta, personal communication and manuscript in preparation). The intracellular uptake of cisplatin, was reported to be mediated by active transporters such as the organic cation transporter 2 (OCT2) and the multidrug toxin extrusion 1 (MATE1) transporter. Passive diffusion through the lipid membrane was further suggested to be crucial for cisplatin uptake but will not be discussed for FY26 as its uptake has been suggested to be mainly an active one (Eljack et al., 2014; Yokoo et al., 2007).

The platinum compound cisplatin is taken through OCT2. The intracellular uptake of oxaliplatin is mediated by OCT2 and OCT3 whereas carboplatin is not a substrate of the organic action transporters (Ciarimboli et al., 2010; Ciarimboli et al., 2005; Yokoo et al., 2007; Yonezawa & Inui, 2011; Yonezawa et al., 2006). The expression of the OCT2 protein levels was rhythmic in kidney with highest and lowest values at ZT06 to ZT10 and at ZT18 to ZT22, respectively. The renal expression of MATE1 did not follow a circadian pattern (Oda et al., 2014). In addition to the role of OCT2, OCT3 and MATE1 for Pt complexes transport, the copper transporter CTR1 was shown to be involved in the intracellular uptake of cisplatin, oxaliplatin and carboplatin as well (Larson et al., 2008; Song et al., 2004). The copper transporter CTR1 is encoded by the gene Slc31a1 which was shown to be rhythmic with highest mRNA expression in rat and mouse liver during the middle of the active span (Almon et al., 2008; Zhang et al., 2014). Based on these data, copper transporter and organic anion transporter might be involved in the intracellular uptake of FY26 as well.
However, the exposure of ovarian cancer cell line A2780 to 100 µM verapamil, an unspecific P-gp inhibitor, and FY26 led to a 1.5-fold accumulation of intracellular concentration of osmium as compared to non-verapamil exposed cells (Coverdale et al., 2018). Thus, there may also be some further role for an active efflux of FY26 or FY25 that could be mediated by P-gp (MDR1 transporter in the ABC transporter family). These transporters also displayed circadian rhythms and were controlled by the molecular clock. The efflux transporter multidrug-resistance genes 1 and 2 (Abcb1 and Abcb4) and the multidrug resistance-associated protein 2 (MRP2) exhibited a circadian rhythm in their mRNA expression in liver, kidney and intestine in male C57BL/6 mice (Ando et al., 2005; Okyar et al., 2012; Zhang et al., 2008). Whereas Abcb1 expression was highest in the middle of the active phase, those of Abcb4 and Mrp2 were in antiphase, with an acrophase in the middle of the inactive phase (Zhang et al., 2008). In a separate work, together with colleagues at Warwick, INSERM, and Istanbul University, I have shown that P-gp mRNA, and protein expressions, as well as function are controlled by the circadian clock both in vitro and in vivo, in mouse ileum, liver and whole organism (Alper Okyar1 et al., 2018a). Thus a circadian rhythm in Abcb1/4 and Mrp2 transporters could also contribute to the chronopharmacokinetics of FY26. Indeed, both transporters were highly expressed near the middle of the inactive phase when mice core body temperature pattern was lowest. This could thus be relevant for FY26 chronopharmacokinetics and chronopharmacodynamics as well. Future experiments could thus aim at the determination of the circadian expression of Abcb1/4 in the time dependent efflux of FY26 in Hepa1-6 Per2-luc cells.

3.2 Chronopharmacodynamics of FY26

Once FY26 was taken up by the Hepa1-6 Per2-luc cells, cells doubling time showed a circadian pattern with the slowest proliferation rate following drug addition at T18. This was confirmed using Per2-luc bioluminescence inhibition as a biomarker. The highest FY26-induced bioluminescence inhibition was observed following drug addition at T22, using the differences between bioluminescence AUCs in drug-exposed vs. controls as an endpoint (AUC
\text{Diff}). I further investigated the role of the circadian clock on the cell cycle phase distribution following FY26 administration at T04 or T16. Exposing Hepa1-6 Per2-luc cells to 7.5 µM FY26 at T04, resulted in an accumulation of cells in G2/M-phase. At the same time no increase of cells in G0/G1-
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phase was observed suggesting that cells did not re-enter the cell cycle, but rather underwent apoptosis. This was not the case for cells exposed at T16. Here, a higher number of cells accumulated in S-phase. A small increase in G2/M-phase cells was also observed, suggesting that fewer cells entered apoptosis at T16 as compared to T04. The induction of apoptosis has been linked to the circadian clock, with overexpression of clock protein PER1 accounting for an increase of apoptosis in radiations-exposed HCT116 cells (Gery et al., 2006). The findings further corroborated the fact that the proliferation of Hepa1-6 Per2-luc cells was slowed down following dosing at T16. Thus, T22 was not the only time point when FY26 addition resulted in best tolerability, using IC50 as an endpoint, but also close to that (T16) when the drug slowed cellular proliferation yet maintained their viability. This was in contrast to T04, when more cell deaths were encountered.

The dose accumulation of Hepa1-6 Per2-luc cells in S-phase that we further found in non-synchronised cell cultures exposed to FY26 was in line with published data in FY26-exposed A549 cells (van Rijt et al., 2014). Such S-phase cells accumulation had also been reported for cisplatin (Velma et al., 2016; Wagner & Karnitz, 2009). On the other hand, following the oxaliplatin treatment of colon cancer (HCT29) or breast cancer cells (MCF7), most cells arrested in G0/G1-phase, whereas the counts of S-phase cells declined (William-Faltaos et al., 2007). Carboplatin produced mixed cell cycle effects pending upon duration of exposure in a bladder cancer cell model: an arrest in S-phase after a 24-h exposure time, as observed for cisplatin, and a decrease in G2/M phase after 48-h (Wang et al., 2010). The FY26-exposed shBmal1-silenced Hepa1-6 Per2-luc cells accumulated in G2/M-phase whilst the number of cells in G0/G1-phase decreased. Additionally, it was observed that the number apoptotic Hepa1-6 Per2-luc shBmal1 was twice as high as compared to Hepa1-6 Per2-luc following the exposure of 7.5 μM FY26. Both observations collaborate the fact that the clock disruptive cell line is more sensitive towards FY26 leading to the determination of a higher number of apoptotic cells.

Whilst the vehicle exposure did not initiate apoptosis, a low induction of apoptosis was observed after the 24 h exposure of IC50 FY26 to Hepa1-6 Per2-luc (15.5% apoptotic cells) and Hepa1-6 Per2-luc shBmal1 cells (7.5% apoptotic cells). The same observation was made for the human ovarian cancer cells (A2780) suggesting that
FY26 did not immediately result in the induction of apoptosis as observed for cisplatin (Henkels & Turchi, 1999; Matsumoto et al., 2016; Romero-Canelón et al., 2015). Besides apoptosis, the induction of autophagy following FY26 exposure showed a trend towards a dose dependent increase in LC3B levels which was higher in the Hepa1-6 Per2-luc shBmal1 cells and immediately after the end of the 24 h FY26 exposure. 

Thus, the results suggest a clock dependent induction of FY26-induced cell death. Indeed, a trend towards the highest expression of the autophagy marker LC3B and a highest number of apoptotic Hepa1-6 Per2-luc shBmal1 cells were encountered as compared to clock-proficient cells. The results, combined with data from the literature, suggested that autophagy and apoptosis could be activated through elevated ROS levels, which has been known to first induce autophagy then apoptosis in case of failure to repair. The induction of apoptosis and autophagy are mediated by increased levels of reactive oxygen levels (ROS) through the inhibition of the autophagy protein Beclin 1 by the anti-apoptotic and protein Bcl-2, which follows a circadian expression with an acrophase at ZT03 to ZT04. (Granda et al., 2005; Higuchi et al., 1998; Pattingre et al., 2005; Redza-Dutordoir & Averill-Bates, 2016; Scherz-Shouval et al., 2007). The peroxide H2O2 inactivates Atg4 which promotes the lipidation of ATG8. This further initiates the formation of the autophagosome leading to the upregulation of LC3B (Barth et al., 2010; Gao et al., 2018; Gouzi et al., 2018; Scherz-Shouval et al., 2007). Thus, circadian pharmacodynamics could be linked to energy metabolism and ROS production, although these were not determined in my thesis. For instance, mitochondrial respiration and electron carrier NAD biosynthesis were rhythmic and clock controlled in mouse liver (Jacobi et al., 2015; Peek et al., 2013; Schmitt et al., 2018).

Unlike the platinum compounds, which are detoxified by reduced glutathione (GSH) (Li et al., 1997), FY26 was reported to be bioactivated by GSH (Fu et al., 2010; Needham et al., 2017). GSH was highest expressed during the mid-active phase in mouse liver (Davies et al., 1983; Xu et al., 2012). High intracellular GSH levels would account for a higher bioactivation of FY26 into FY25. As the highest concentrations of GSH in mouse tissues occurred in liver, and the acrophase of this rhythm was located at ZT18-ZT22 (Li et al. 1997), one can expect highest bioactivation of FY26
to occur in the second half of the nocturnal activity span of mice. Additionally, a high bioactivation rate would thus account for a high efficacy at ZT18 which was confirmed by the chronoefficacy study as the highest tumour growth inhibition occurred at ZT18. Nevertheless, further experiments investigating the chronoefficacy will be needed, as this study was done involving a small number of mice. Moreover, the determination of circadian GSH levels together with ROS levels, following the exposure of FY26 in Hepa1-6 Per2-luc cells, are crucial because GSH is involved in the detoxification of ROS. GSH decreases ROS levels (Armstrong et al., 2002; Tan et al., 1998). As GSH expression is highest during the mid-active span in mice, low ROS levels would have been expected. In this context, the determination of circadian GSH and ROS levels following FY26 exposure will allow to track the pharmacokinetics-pharmacodynamics interactions in Hepa1-6 Per2-luc cells and connect this to in vivo chronotoxicity and chronoefficacy.

Unfortunately, the measurement of ROS was not possible as FY26 interaction with ROS using the Seahorse instrument. This methods will be replaced in future experiments by trying alternative ROS measurements by using DCFDA (2’-7’-diclorfluorescein).

### 4 FY26 and its interaction with the circadian clock

FY26 effects on the circadian clocks were assessed using circadian clock gene Per2 reporter expression both in cell cultures and in mice, clock genes mRNA circadian expressions in cell cultures, and core body temperature circadian rhythm in mice.

In temperature cycle synchronised Hepa1-6 Per2-luc cells, FY26 transiently reduced bioluminescence, yet circadian oscillations persisted. The amplitude of the circadian oscillations were largest following FY26 dosing at T22. The persistence of the Per2-luc circadian oscillations, despite FY26 exposure, suggested that the circadian clock could remain functional. This finding was further supported by the persistent oscillatory patterns in the mRNA expression of clock genes Per2, Bmal1 and Rev-erba, as determined with RT-qPCR in similar experimental conditions. This suggest that the decreased amplitude in Per2-luc bioluminescence was not caused by the interaction of FY26 with Per2 but by FY26 cytotoxicity instead.
In contrast to FY26, cisplatin increased the mRNA expression of both Per2 and clock controlled gene Dbp, whilst abolishing Cry2 and Rev-erba rhythms in the liver of C57BL/6 mice (Cao et al., 2018). The cytokine interferon-α dampened the Per2 rhythm in ICR mouse liver with the largest amplitude reduction following dosing at ZT14 (Ohdo et al., 2001; Shinohara et al., 2008). Topoisomerase I inhibitor irinotecan down regulated and suppressed circadian mRNA expressions of clock genes Rev-erba, Per2 and Bmal1 in the liver of male or female B6D2F1 mice, with mild or severe clock disruption depending upon dosing time (Ahowesso et al., 2011). Antimetabolite 5-fluorouracil reduced the expression of Per1 and Per2 in mouse liver as well as in NIH3T3 fibroblast without affecting cell viability (Terazono et al., 2008). FY26 effected the CTS at variance with a strong clock disruption induced by the above-mentioned anticancer drugs. In the hepatocarcinoma model, FY26 did not affect the clock gene mRNA expression, but altered cell cycle and cell death mechanisms. Thus, the time dependent decrease in Per2-luc bioluminescence amplitude were considered as a reflection of FY26-induced transient arrest of cell proliferation, rather than drug interaction with the Per2 promoter in Hepal-6 Per2-luc cells. However, FY26 interacted with PER2 protein measured through PER2::LUC liver bioluminescence. Following a single i.p. injection of FY26 led to the ablation of PER2::LUC rhythmicity which was highest at CT06, referring to CT22.5 to CT01.5, as the AUC of bioluminescence was lowest accounting for the highest bioluminescence inhibition. This was thus in alignment with the highest in vivo FY26 toxicity at ZT18.

The effect of FY26 on core body temperature was simultaneously measured with that on PER2 protein expression. At both circadian time points, FY26 injection caused a drop in core body temperature. At the CT24, the FY26 injection led to an ablation of rhythmicity which recovered 2 days after the injection. However, injecting FY26 at CT12, led to an increase of the amplitude of core body temperature rhythms followed by a recovery 2 days after the injection. The same experiment was further conducted on mice kept under constant LD 12:12 entrainment. Here, the single i.p. injection of FY26 led to a sharp decrease of core body temperature at ZT18 whereas no such effect was observed following the injection at ZT06. This further confirmed the previously identified time point of worst FY26 tolerability in male C57BL/6 mice. Several other cancer drugs were found to affect core body temperature rhythm. Irinotecan disrupted
core body temperature rhythm mostly following its administration at the least tolerated time point ZT03 in female mice. Less alterations of daily circadian body temperature rhythm was found for male mice receiving irinotecan (Ahowesso et al., 2011). Interferon administration at ZT12 dampened the amplitude of rectal temperature more as compared to ZT0 (Ohdo et al., 2001). A further drug which effects core body temperature when given at the time point of least tolerability, is the antimetabolite gemcitabine (Li et al., 2005a). Thus, FY26 like other anticancer drugs altered the circadian rhythm in core body temperature predominantly following its administration at the time point of least tolerability.

Besides, the circadian disruption of core body temperature, the data further pointed towards the crucial role of circadian entrainment with regard to toxicity. FY26 affected core body temperature rhythm when injected at CT24 and CT12 in constant darkness. However, no such effect was observed when FY26 injected at ZT06 in LD 12:12 synchronised mice. As mentioned in the first chapter of the discussion, the accumulation of Per2 and Hsp in clock disruptive mice can result in circadian rhythms independently from the clock (Kornmann et al., 2007). This might as well suggests that PER2 plays a crucial role in the maintenance of core body temperature rhythms as an interaction with PER2 could lead to disruptive core body temperature rhythms. The extent to which core body temperature rhythm was disrupted, appeared to depend on whether the mouse was kept under alternating LD 12:12 entrainment or constant darkness. As a reduced toxicity was observed in mice kept under LD 12:12 entrainment, light-dark synchronisation appeared to be crucial to coordinate a circadian pattern together with lowering FY26 toxicity. Translating this to the chronotherapy of cancer patients, would mean that maintaining a stable daily routine is important as it could contribute to the reduction of the toxicity of chemotherapy. As mentioned in the introduction (chapter I.9), the disruption of the CTS was associated with a lower survival rate in metastatic breast and gastro-intestinal cancer patients (Ballesta et al., 2017; Sephton et al., 2000). This was further confirmed by the effect of FY26 on core body temperature. Thus, developing anticancer drugs which do not affect the CTS could indeed help increase survival and should thoroughly be considered in the development of anticancer agents.
X Conclusion
The in vitro and in vivo studies were designed to answer to the questions (i) whether FY26 displays a circadian variation in its pharmacokinetics and pharmacodynamics characteristics and (ii) whether endogenous temperature cycles can be used to transfer in vitro chronotoxicity into in vivo chronotoxicity.

I have demonstrated that the tolerability of FY26 was best following dosing during the first half of the inactive phase (ZT06) in mice and near the nadir of rhythmic temperature exposure in cell cultures (T22). FY26 chronotolerance pattern in cell cultures was aligned with that in mice, using temperature cycle as a biomarker. Indeed, FY26 tolerability was best, both in vitro and in vivo, following its administration near the acrophase of temperature exposure in both experimental models. It is the first time that such link between in vitro and in vivo chronotoxicity is shown, to the best of my knowledge. I have further discussed that time dependent pharmacokinetics and pharmacodynamics, including detoxification, of FY26 might result in part from the circadian expression of efflux transporters Mdr1 or Mrp2. Interestingly intracellular FY26 uptake was highest at T02 whereas the minimum toxicity occurred at T22. This was consistent as a higher number of cells entered apoptosis at T04 whereas at T16 a higher number of cells accumulates in S-phase thus decelerate cell proliferation and bioluminescence.

FY26 did not seem to alter the molecular circadian clock, at the level of mRNA gene expression, but rather at protein expression levels in cell populations. In mice (whole organism), a marked alteration of the core body temperature rhythm was uncovered. The intensity of the effect of FY26 on the protein expression and on core body temperature varied as a function of whether the mice were kept in LD12:12 synchronisation or under constant darkness. The administration of FY26 at CT24 and CT12 led to a disruption of PER2 protein expression at CT12 whereas at CT24 the amplitude of the oscillation was dampened.

Regarding core body temperature, FY26 disrupted the core body temperature circadian rhythm in 2 out of 3 mice at both time points tested that corresponded to endogenous mean circadian times CT24 and to CT12. Neither time points matched the expected CTs of best and worst tolerability because of the free-running conditions in DD. In mice, kept under LD 12:12 entrainment, core body temperature rhythm was not altered after FY26 dosing at ZT06. In contrast, core body temperature dropped by
3.7°C then recovered within 24 h following dosing at ZT18. Thus, FY26-induced circadian disruption was worse following dosing at ZT18, using both PER2::LUC bioluminescence and circadian temperature rhythms as endpoints, in agreement with the chronxicity pattern of FY26.

Contrary to other anticancer agents, FY26 tended to display the highest tumour inhibition properties at ZT18. However, as the tumour efficacy studies could not be repeated, further research is needed to delineate the chronoefficacy of FY26.

Further limitation to our study involved the humane endpoints of the in vivo studies. Once a body weight loss of 20% of pre-treatment body weight was recognised, the mouse was euthanized regardless of any sign of pain. As body weight was chosen as an endpoint for most in vivo studies described in this thesis and those reported in the literature, a maximum allowance of 20% body weight loss only enabled the detection of larger variations of body weight loss after FY26 administration at different circadian time points. This led to the early exclusion from the study of mice losing weight despite being clinical well. Regarding the in vitro studies, the experiments highlight the need for a systematic mapping of the circadian mechanisms of action of FY26, including time dependent active uptake and possibly ABCB4 and MRP2-mediated efflux, the implication of circadian changes in GSH levels for both bioactivation of FY26, and its detoxification, and the time dependent effect of FY26 on ROS induction. Such mechanisms will lend support to the chronopharmacokinetics and chronopharmacodynamicics that have been demonstrated here in synchronised cell culture models.

The investigation of the chronopharmacology of FY26 and the determination of the effect of FY26 on the circadian clock are crucial. Clock disruption results in the impairment of cancer therapy through the promotion of tumour development and growth. (Innominato et al., 2012; Mormont et al., 2000; Sulli et al., 2018). Therefore, developing a drug as FY26 which was shown to overcome cisplatin resistance, to exhibit a circadian toxicity and efficacy profile with minimal clock interaction when given at ZT06, lowers adverse effects of chemotherapy as well as it enhances tumour growth inhibition. So far, FY26 is understood as a novel, potential anticancer drug to overcome cisplatin resistance. The use of endogenous temperature cycles identified the circadian in vitro tolerability profile of FY26 but also allowed predictions on the
In conclusion, the in vivo chronotoxicity of FY26. Thus, temperature cycles helped to unravel the FY26 chronopharmacology whilst it has the potential to minimise the number of mice used in in vivo experiments.

In order to put this into a larger context, the use of temperature cycles mimicking core body temperature rhythm as marker of the CTS may constitute a crucial new factor in drug development. CTS robustness has been linked to significant improvements in both patient’s survival and quality of life in cancer patients. We show that the information it conveys could also help reduce animal experiments, an idea which is in alignment with the 3R’s of animal research (replacement, reduction and refinement) and supported by the public opinion on the use of animals for research.
XI Appendices
1 Appendices: *In vitro* chronotoxicity

![Dose response curve of Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 under the exposure of the concentration scale A.](image)

*Figure XI-1*: Dose response curve of Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 under the exposure of the concentration scale A.

Four thousand Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 cells were seeded in each well of a 96-well NUNC plate. Two days after cell seeding, cells were exposed to different FY26 concentrations (0.001 µM, 0.01 µM, 0.1 µM, 1 µM, 10 µM, 100 µM FY26) of the concentration scale A. After 24 h of incubation, FY26 was removed and replaced by media. The total cellular protein content was measured using SRB-Assay (n=12) 72 h after FY26 withdrawal. The cell survival was calculated. For Hepa1-6 Per2-luc the maximum cell survival was 98.0% ± 0.6% at 0.001 µM FY26 and the minimum was 6.9% ± 0.2% for 100 µM FY26. The highest survival rate of Hepa1-6 Per2-luc shBmal1 was 109.5% ± 2.6% following exposure to 0.001 µM and the lowest one was 6.4% ± 0.3% for 100 µM.
Figure XI-2: Optimisation of the concentration scales done in Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1.

Four thousand Hepa1-6 Per2-luc and Hepa1-6 Per2-luc shBmal1 cells were seeded per well of a 96-well NUNC plate and exposed to different FY26 concentrations after 2 days (n=12). Cells were exposed for 24 h to concentration scales B, C and D. Total cellular protein content was measured 72 h after FY26 removal. For both, Hepa1-6 Per2-luc and its shBmal1 clone, a maximum cell survival of 101.2 ± 1.2% and 101.2% ± 0.2% and a minimum cell survival of 4.2% ± 0.2% and 4.2% ± 0.1% were achieved. Concentration scale C was used for exposing Hepa1-6 Per2-luc resulting in a survival rate ranging from 21.2% ± 1.5% to 97.5% ± 1.5%. Further, a minimum and maximum survival of 16.3% ± 2.1% to 100.8% ± 2.0% was observed Hepa1-6 Per2-luc shBmal1 under the exposure of the concentration scale D.
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Figure XI-3: Optimisation of the concentration scales in HCT116 Per2-luc.

Eighty-thousand cells have been seeded and drugged with different concentrations of FY26. Experiment 1 and 2 were exposed to the concentration scale H and experiment 3 to the concentration scale I for 24 h. Total cellular protein content was measured 72 h after the drug removal. One-way ANOVA showed a significant difference between the IC_{50} values with p<0.0001. Tukey’ multiple comparison detected a direct significant difference between the IC_{50} of experiment 1 vs. C (****), experiment 2 vs. 3 (****) as well as between experiment 1 vs. 2 (**). The cell survival for the 1st experiment ranged from 6.8% ± 0.5% to 89.9% ± 2.9%, for the 2nd experiment from 49.4% ± 9.1% to 97.2% ± 1.2% and for the third experiment from 4.7% ± 0.4% to 99.9% ± 0.2%.
Figure XI-4: Dose response curves at T04, T10, T16 and T22 in Hepa1-6 Per2-luc after the exposure of the concentration scale E (Experiment A).

Different concentrations of FY26 of the concentration scale E (16 µM, 12 µM, 10 µM, 8 µM, 6 µM, 4 µM, 2 µM, 1 µM and 0.5 µM) were exposed at T04, T10, T16 and T22 after 2.5 days after the start of temperature synchronisation. After 24 h exposure time, the drug-media was replaced by fresh media. The total protein content was measured 72h after media change (n=18). Minimum and maximum cell survival rates (mean ± SEM) ranged from 3.9% ± 0.1% to 89.5% ± 2.2% for T04, from 3.5% ± 0.1% to 93.1 ± 1.6% for T10, from 3.6% ± 0.1% to 92.1% ± 1.9% for T16 and from 3.4% ± 0.1% to 90.7% ± 2.2% for T22 using concentration scale E.
Figure XI-5: Dose response curves at T04, T10, T16 and T22 in Hepa1-6 Per2-Luc after the exposure of the concentration scale F (Experiment B).

Different concentrations of FY26, concentration scale F, (30 µM, 15 µM, 10 µM, 8 µM, 6 µM, 4 µM, 2 µM, 1 µM and 0.5 µM) were used for exposing synchronised cell cultures at T04, T10, T16 or T22 after 2.5 days after the start of temperature synchronisation. After 24 h exposure duration, the drug-containing media was replaced by fresh media. Total protein content was measured 72h after media change (n=12) using the SRB assay. The minimum and maximum survival rate of cells subjected to concentration scale F was 7.9% ± 0.4% to 99.9% ± 0.2% for T04, 11.9% ± 0.5% to 99.9% ± 0.4% for T10, 7.6% ± 0.6% to 100.5% ± 0.15 for T16 and 10.4% ± 0.4% to 98.8% ± 0.2% for T22.
After 2.5 days of synchronisation of Hepa1-6 Per2-luc shBmal1 cells were treated with 21 $\mu$M, 14 $\mu$M, 7 $\mu$M, 6 $\mu$M, 5 $\mu$M, 4 $\mu$M, 2 $\mu$M, 1 $\mu$M and 0.5 $\mu$M. The media was changed after 24 h drug exposure time. Protein content was determined 72 h after media change ($n=12$). The minimum survival ranged from 26.0% ± 2.3% to 99.5% ± 0.5% for T04, from 21.3% ± 1.6% to 100.6% ± 0.3% for T10, from 19.3% ± 1.3% to 99.7% ± 0.3% for T16 and from 25.8% ± 1.8% to 100.2% ± 0.2% for T22.

Figure XI-6: Dose response curve of Hepa1-6 Per2-luc shBmal1 at T04, T10, T16 and T22 after the exposure of the concentration scale $G$. 
Figure XI-7: Dose response curve at T04, T10, T16 and T22 for Hepa1-6 Per2-luc shBmal1 after the exposure of the concentration scale F.

Hepa1-6 Per2-luc shBmal1 cells were incubated for 24 h at different FY26 concentrations (30 µM, 15 µM, 10 µM, 8 µM, 6 µM, 4 µM, 2 µM, 1 µM and 0.5 µM). Media changed was done 24 h after drug administration and protein content was determined 72 h after media change (n=12). A minimum to maximum survival was obtained ranging from 15.4% ± 1.2% to 99.6% ± 0.1% for T04, from 13.0% ± 1.1% to 99.8% ± 0.1% for T10, from 11.9% ± 0.3% to 99.9% ± 0.4% for T16 and from 22.9% ± 3.9% to 99.0% ± 0.2% for T22.
Figure XI-8: Dose response curves at T04, T10, T16 and T22 on synchronised HCT116 Per2-luc cells after exposure of the concentration scale I.

Eighty thousand HCT116 Per2-luc cells were seeded per well in a 96-well FALCON plate and subjected to temperature cycle D. Different concentrations of FY26 were added after 2.5 days of synchronisation at T04, T10, T16 or T22. FY26-containing media was removed and replaced with drug-free media after an exposure duration of 24 h. Cellular protein content was determined in each dish 72 h after completion of drug exposure (n=23). Solid and dotted lines link experimental dose-related survival rates in first and second experiment respectively. The minimum cell survival ranged from 7.9% ± 1.7% to 11.7% ± 2.2% and the maximum survival varied from 94.6% ± 1.7% to 104.7% ± 2.2%.
2 Appendices: *In vivo* Chronotoxicity

**Figure XI-9:** Escalating dose selection study for FY25 and FY26.

Panel A and B show the body weight development of non treated mice, mice treated with the vehicle, 20, 40, 80 and 160 mg/kg FY25 (panel A) and FY26 (panel B). Panel C depicts the survival of mice treated with FY25 (Log-rank test $p=0.0002$) and panel D shows the survival of mice treated with FY26 (Log-rank test $p<0.0001$).
Figure XI-10: Histopathological slices of mouse stomach.

Mouse 1 was injected with the vehicle (control) and mouse 2 to mouse 5 were injected with 50 mg/kg FY26.
Figure XI-11: Histopathological slices of mouse ileum.

Mouse 1 was injected with the vehicle (control) and mouse 2 to mouse 5 were injected with 50 mg/kg FY26.
Figure XI-12: Histopathological slices of mouse colon.

Mouse 1 was injected with the vehicle (control) and mouse 2 to mouse 5 were injected with 50 mg/kg FY26.
Figure XI-13: Histopathologic slices of mouse liver.

Mouse 1 was injected with the vehicle (control) and mouse 2 to mouse 5 were injected with 50 mg/kg FY26.
Figure XI-14: Histopathological slices of mouse kidney.

*Mouse 1 was injected with the vehicle (control) and mouse 2 to mouse 5 were injected with 50 mg/kg FY26.*
Figure XI-15: Histopathological slices of mouse spleen.

Mouse 1 was injected with the vehicle (control) and mouse 2 to mouse 5 were injected with 50 mg/kg FY26.
Figure XI-16: Histopathological slices of mouse bone marrow.

Mouse 1 was injected with the vehicle (control) and mouse 2 to mouse 5 were injected with 50 mg/kg FY26.
Figure XI-17: In vitro and in vivo temperature cycles recorded with the Anipill®.

The figure depicts the temperature cycles measured by the Anipill® which were placed in three 35 mm NUNC Petri dishes put in the cell culture incubator and implanted in the peritoneal of three male C57BL/6 mice.
3 Appendices: FY26 and the circadian timing system

3.1 Result of Spectrum Analysis on mice core body temperature under LD 12:12 synchronisation
Figure XI-18: Changes of mice core body temperature after the injection of the vehicle at ZT06 and ZT18. The black marker represents the i.p. injection of the vehicle.
Figure XI-19: Changes of mice core body temperature rhythm after the injection of 50 mg/kg FY26 at ZT06 and ZT18. The black marker represents the i.p. injection of 50 mg/kg FY26. The vertical line marked the injection time.
3.2 Results of the Spectrum Analysis on the PER2::LUC oscillation
CT06 Vehicle mouse 7 – normalised photon counts

CT06 FY26 mouse 1 – normalised photon counts
Figure XI-20: Results of the Spectrum Analysis showing the changes of the PER2::LUC expression and its period, acrophase and amplitude in 9 PER2::LUC mice after the treatment with the vehicle and 50 mg/kg FY26 at CT06 and CT18.

The vertical line marked the injection time point. Mouse 1 to 3 were reallocated in the RT-Bio on the 11/12/17 (not shown in graph), mouse 4 to mouse 6 were reallocated on the 05/02/18 and mouse 7 to mouse 9 on the 22/02/18.
3.3 Spectrum Analysis of mice core body temperature in without external synchroniser (DD)

Core temperature CT06 FY26 mouse 4 (RT-Bio)

Core temperature CT18 FY26 mouse 5 (RT-Bio)
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Core temperature CT18 Vehicle mouse 6 (RF-Bio)

Core temperature CT06 Vehicle mouse 7 (RF-Bio)
Figure XI-21: Results of the Spectrum Analysis on core body temperature of 9 PER2::LUC mice when injected with the vehicle and 50 mg/kg FY26 at CT06 and CT18.

Mouse 1 to 3 were reallocated in the RT-Bio on the 11/12/17 (not shown in graph), mouse 4 to mouse 6 were reallocated on the 05/02/18 and mouse 7 to mouse 9 on the 22/02/18. The vertical line marked the injection time.
XII Bibliography


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