Anti-tumour compounds illudin S and Irofulven induce DNA lesions ignored by global repair and exclusively processed by transcription- and replication-coupled repair pathways

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Abstract

Illudin S is a natural sesquiterpene drug with strong anti-tumour activity. Inside cells, unstable active metabolites of illudin cause the formation of as yet poorly characterised DNA lesions. In order to identify factors involved in their repair, we have performed a detailed genetic survey of repair-defective mutants for responses to the drug. We show that 90% of illudin’s lethal effects in human fibroblasts can be prevented by an active nucleotide excision repair (NER) system. Core NER enzymes XPA, XPF, and TFIIH are essential for recovery. However, the presence of global NER initiators XPC, HR23A/HR23B and XPE is not required, whereas survival, repair and recovery from transcription inhibition critically depend on CSA, CSB and UVS, the factors specific for transcription-coupled NER. Base excision repair and non-homologous end-joining of DNA breaks do not play a major role in the processing of illudin lesions. However, active RAD18 is required for optimal cell survival, indicating that the lesions also block replication forks, eliciting post-replication-repair-like responses. However, the translesion-polymerase DNA pol y is not involved.

We conclude that illudin-induced lesions are exceptional in that they appear to be ignored by all of the known global repair systems, and can only be repaired when trapped in stalled replication or transcription complexes. We show that the semisynthetic illudin derivative hydroxymethylacylfulvene (HMAF; Irofulven), currently under clinical trial for anti-tumour therapy, acts via the same mechanism.

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Abbreviations: BER, base excision repair; CS, Cockayne syndrome; GG-NER, global-genome nucleotide excision repair; HMAF, hydroxymethylacylfulvene; NHEJ, non-homologous end-joining; PRR, post-replication repair; TC-NER, transcription-coupled nucleotide excision repair; UDS, unscheduled DNA synthesis; UVS, UV-sensitivity syndrome; XP, xeroderma pigmentosum

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1. Introduction

The illudins are a family of natural sesquiterpene compounds with anti-tumour activity, originally isolated from the mushrooms *Omphalotus illudens* (jack-o-lantern) or *Lampteromyces japonicus* [1]. Although many illudins are highly effective against various drug-resistant tumours both in vivo and in vitro, the extreme cytotoxicity of these compounds in the nanomolar range has severely restricted their practical use in cancer therapy [2,3]. Recently semisynthetic derivatives have been reported [4] with a strongly improved therapeutic index [2,5,6] of which hydroxymethylacylfulvene (HMAF, Irofulven) is currently under clinical trial [7,8]. The tissue specificity and tumour selectivity of the illudins has been attributed to the presence of an energy-dependent system mediating transport into the cells [9] and subsequent metabolic activation to an unknown reactive intermediate [10]. Inside the cells, strong inhibition of DNA synthesis occurs, presumably caused by DNA damage [2]. The requirement for prolonged incubation periods of actively metabolising cells has hampered the identification of the responsible DNA adduct [11].

Table 1
Properties of cells strains used

<table>
<thead>
<tr>
<th>Human fibroblast strain</th>
<th>Clinical symptoms</th>
<th>Affected gene</th>
<th>NER activity (%)</th>
<th>Repair pathway affected</th>
</tr>
</thead>
<tbody>
<tr>
<td>CSRO</td>
<td>Normal control</td>
<td>–</td>
<td>100 (deficient)</td>
<td>–</td>
</tr>
<tr>
<td>XP25RO</td>
<td>XP with neurological complications</td>
<td>XPA</td>
<td>&lt;1</td>
<td>GG-NER + TC-NER</td>
</tr>
<tr>
<td>XPCS1BA</td>
<td>XPCS complex</td>
<td>XFB (ERCC3)</td>
<td>5</td>
<td></td>
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<tr>
<td>XP28RE</td>
<td>XP with neurological complications</td>
<td>XPD (ERCC2)</td>
<td>25</td>
<td></td>
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<tr>
<td>XP24RO</td>
<td>Mild XP</td>
<td>XPF (ERCC4)</td>
<td>15–30</td>
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<td>XP28</td>
<td>XP</td>
<td>XPG (ERCC5)</td>
<td>2</td>
<td></td>
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<td>XP21RO</td>
<td>XP</td>
<td>17</td>
<td>GG-NER</td>
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<td>XP31RO</td>
<td></td>
<td>XPC</td>
<td>15</td>
<td></td>
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<tr>
<td>XP24RE</td>
<td></td>
<td></td>
<td>20</td>
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<td>XP27RO</td>
<td>Mild XP</td>
<td>XPE (DDR2)</td>
<td>61</td>
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<tr>
<td>CS5RE</td>
<td>Cockayne syndrome</td>
<td>CSA (ERCC6)</td>
<td>~100</td>
<td>TC-NER</td>
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<td>CS1AN</td>
<td>Cockayne syndrome</td>
<td>CSB (ERCC6)</td>
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<tr>
<td>Kps3</td>
<td>UVS syndrome</td>
<td>Not cloned</td>
<td></td>
<td></td>
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<tr>
<td>XP1RO</td>
<td>Mild XP (variant)</td>
<td>RAD50A (POLH)</td>
<td>~100</td>
<td>Translation DNA synthesis</td>
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<tr>
<td>XP3RO</td>
<td>Non-photosensitive trichothiodystrophy</td>
<td>Not cloned</td>
<td>~100</td>
<td>NER = normal</td>
</tr>
</tbody>
</table>

Other cell strains

<table>
<thead>
<tr>
<th>Species (origin)</th>
<th>Affected gene(s)</th>
<th>Repair pathway affected</th>
</tr>
</thead>
<tbody>
<tr>
<td>43-3B Chinese hamster ovary (CHO9)</td>
<td>ERCC1</td>
<td>NER and crosstalk repair</td>
</tr>
<tr>
<td>EM-9 Chinese hamster ovary (AA8)</td>
<td>XRC1</td>
<td>Base excision repair</td>
</tr>
<tr>
<td>V15B Chinese hamster ovary (Y79B)</td>
<td>Kab6</td>
<td>Non-homologous end-joining</td>
</tr>
<tr>
<td>XR-1 Chinese hamster ovary (CHO-K1)</td>
<td>XRC4</td>
<td></td>
</tr>
<tr>
<td>HR23DKO Double-targeted mouse mutant embryonic fibroblasts</td>
<td>HR23A and HR23B</td>
<td>GG-NER</td>
</tr>
<tr>
<td>XPC– Targeted mouse embryonic fibroblasts</td>
<td>XPC</td>
<td></td>
</tr>
<tr>
<td>DT40 (Aral18) Chicken lymphoma (DT40)</td>
<td>RAD38</td>
<td>Post-replication repair</td>
</tr>
</tbody>
</table>

* Measured as UV-induced unscheduled DNA synthesis (UDS).

a GG, global-genome; TC, transcription-coupled; NER, nucleotide excision repair.
genetic studies using UV-sensitive rodent mutant cell lines have indicated that nucleotide excision repair (NER) is involved in the removal of illudin-induced lesions [11]. However, contribution of other repair pathways such as base excision repair (BER) or recombination repair cannot be excluded as long as the spectrum of lesions remains undefined.

In order to further investigate the mechanisms of the extreme toxicity of the illudins, we have studied the toxicity and repair of illudin S lesions in a variety of mammalian cell strains with known defects in DNA repair processes, including NER, BER, non-homologous end-joining (NHEJ) and post-replication repair (PRR), (listed in Table 1). Surprisingly, the data indicate that illudin-induced DNA adducts are ignored by the global-genome repair pathways (NER, BER and NHEJ), and can only be processed when trapped in sites of stalled transcription or replication forks. Transcription-coupled NER and post-replication repair are the key processes protecting cells to illudin S toxicity.

2. Materials and methods

2.1. Cell strains and culture conditions

Human fibroblast strains were cultured in Ham F10 supplemented with 15% foetal bovine serum and antibiotics. For Chinese hamster ovary cells a 1:1 mixture of Ham F10 and DMEM was used. Chicken lymphoma cells were cultured at 39°C in HEPES-buffered RPMI1640, supplemented with 10% inactivated calf serum and 1% chicken serum. Properties of the strains used in this study are summarised in Table 1.

2.2. Agents

UV light of predominantly 254 nm was emitted from a Philips TUV germicidal tube (fluence rate 0.3 W/m²). Stock solutions of illudin S (0.2 μg/ml corresponding to 0.76 μM), isolated from O. illudens as described [12] and HMAF (Irofulven, 1 mg/ml) are fully stable at 4°C [2] and diluted in culture medium to the desired concentrations, immediately prior to use. Radioactive nucleic acid precursors were all purchased from Amersham Radiochemicals.

2.3. Cell survival assays

To avoid the time-consuming and laborious colony assays of primary human fibroblasts, a simplified approach was followed for measurement of cytotoxicity: Sparse cultures (5 × 10³ to 10 × 10³ cells per 6 cm-dish, or 1 × 10³ to 2 × 10³ per 3 cm-dish) were grown for 48 h and then exposed to illudin S or UV and new medium was added. After 3-5 days of further culture, when untreated control cultures were still far from confluence, cells were pulse-labelled with [methyl-3H]-thymidine (40-60 Ci/mmol, 5 μCi/ml) for 3 h in presence of 20 mM HEPES pH 7.3, to obtain a measure for the number of proliferating cells in each culture. After 30 min chase in unlabelled medium, cells were rinsed, lysed in 0.25 M NaOH and harvested. The incorporated radioactivity, measured in a liquid scintillation counter, was processed into mean ± S.E. points. As reported earlier for human fibroblasts [13,14] as well as rodent cells [11,15], this simplified protocol produces highly reproducible UV-survival curves covering three logs, and indistinguishable from those obtained with classical colony assays. With a few standard strains this was confirmed for illudin. Almost all presented panels with survival curves are from single representative experiments, containing negative and positive controls to ensure reproducibility. Error bars represent variability from triplicate of duplicate dishes.

Chinese hamster cells were sparsely seeded in dishes, exposed to various concentrations of illudin and allowed to form colonies. Chicken lymphoma cells were seeded at 10⁶ cells per ml, exposed to illudin for 24-48 h and counted or checked for thymidine incorporation as above, immediately or after 1 day.

2.4. Unscheduled DNA synthesis

Coverslip cultures were exposed to 16 J/m² of UV light and cultured for 2 h in tritiated thymidine as described [16]. In case of illudin S, cells were incubated with 60 ng/ml in presence of [methyl-3H]-thymidine (120 Ci/mmol, 5 μCi/ml) for 24 h. After radioactive labelling, all cultures were fixed and processed for autoradiography as described.
[16], and silver grains were counted over at least 50 non-S-phase cells.

2.5. RNA synthesis inhibition and recovery

One-day-old fibroblast cultures in 3 cm-dishes were prelabelled with $[^{2-}{^{14}}C]$-thymidine (50 mCi/mmol, 50 nCi/ml) for 16 h, exposed to illudin S for 3 h and cultured further. At various times, cells were pulse-labelled (1 h) with $[^{5,6-}{^{3}}H]$-uridine (38 Ci/mmol, 2 μCi/ml), alkali-lysed and processed for scintillation counting as described above. The ratios of $^{3}H$ to $^{14}C$ are taken as a measure of overall RNA synthesis [13].

3. Results

3.1. Core NER factors are required for repair of illudin damage

Based on earlier experience with SV40-transformed human fibroblasts [9] a treatment protocol of 72 h with doses up to 1 ng/ml (0.26 nM) of illudin S was chosen for the primary fibroblasts. In these conditions, survival decreased exponentially with dose, revealing a $D_{10}$ value of about 1.1 ng/ml in normal control cells and 0.12 ng/ml for fully NER-deficient XP-A fibroblasts (see Fig. 1A). Thus, normal cytotoxicity was slightly higher than observed in a single
SV40-transformed normal human fibroblast culture [9]. The XP-A hypersensitivity ratio of 8–10 was about the same as that obtained for UV-exposure, with $D_{10}$ values of about 1 and 10 J/m² for XP-A and normal cells, respectively (Fig. 1B). With shorter exposure times, used in later experiments, higher doses were required to produce the same lethality. An inverse relationship between time and dose was found to exist between concentrations of 0.5 and 10 ng/ml at least, with no significant change in relative sensitivity of NER-defective XP-cells (Figs. 1A and C and 3A and B, and data not shown).

In UV-sensitive fibroblasts with defective XPB, XPD, XPF and XPG, other core enzymes of the NER-pathway, the responses to illudin S were similarly exaggerated, though less pronounced. In all these cases, the hypersensitivity factors after exposure to illudin S and UV irradiation were comparable (Fig. 1A, B and D), confirming the requirement of NER for removal of illudin lesions, and suggesting similar mechanisms of cytotoxicity.

3.2. Global-genome NER is dispensable

Unexpectedly, cells from XP group C, presented a clear exception: an intermediate sensitivity to UV (2.5–5) times (Fig. 1B), typical of XP-C strains [17] contrasted with a fully normal survival after illudin S exposure, a response found in three different human XP-C strains (Fig. 2A). In XPC-defective mouse fibroblasts, known to be sensitive to UV as well [18], this behaviour was confirmed (Fig. 2D). The possibility existed that normal XP-C survival was simply related to the protracted low dose treatment protocol in combination with residual repair activity. This explanation is highly unlikely since moderately UV-sensitive XP-F cells with a residual NER level of 15–30% (similar to XP-C), were as sensitive to illudin as they were to UV (Fig. 1D). In order to rigorously exclude potential dosimetry effects, we confirmed XP-C normalcy in a shorter (24 h) treatment protocol (Fig. 2B). Furthermore, both XP-A and XP-C cells retained their sensitivity to UV, when irradiation was split in seven doses delivered over a 72 h period, in order to mimic a low-dose-rate regimen similar to illudin S (Fig. 2C). Consistent with the unexpected behaviour of XP-C cells, UV-sensitive mouse fibroblasts defective in both HR23A and HR23B proteins, which can bind XPC protein and stabilise it (Ng et al., manuscript in preparation), were also as resistant to illudin as normal mouse cells (Fig. 2D). In addition to XP-C cells, also XP-E cells survived illudin exposure normally (see Fig. 1D).

3.3. Illudin lesions are exclusively removed by transcription-coupled NER

The enzyme complexes containing XPC and XPE are responsible for the lesion-recognising steps in the NER subpathway which repairs global DNA damage. In XPE and XPC mutants preferential repair of actively transcribed DNA strands is fully intact. Consistent deficiency (to varying degrees) in both global-genome repair (GG-NER) and transcription-coupled repair (TC-NER) is a characteristic feature of the other XP groups (for a review, see [19]). Our data suggest that only TC-NER is relevant to illudin cytotoxicity. Therefore, cells from CS patients, known to have a selective defect in TC-NER, were studied. CS-A and CS-B cells were seven and nine times more sensitive to illudin than two cell strains with normal NER (see Fig. 3A). These responses to illudin are much more pronounced than to UV (two and four times, respectively, data not shown here. See [13,17]) and more near to fully NER-deficient XP-A cells (see Fig. 3C). We found exactly the same hypersensitivity pattern in cells from patients with UV-sensitivity syndrome (UVS, [20,21] see Fig. 3C), which also suffer from defective TC-NER [22,23].

TC-NER is initiated at transcription forks stalled at DNA lesions and allows completion of the transcript after removal of the lesion (for overview see: [24]). To study the inhibition and recovery of RNA synthesis, the fibroblasts were exposed to illudin in a shortened, 3 h-exposure protocol, followed by a pulse-labelling with tritiated uridine at various times (Fig. 3B). Initial inhibition and subsequent recovery were evident in normal and XP-C cells, but did not at all occur in fully NER-defective XP group A. We conclude, that illudin S-induced lesions indeed inhibit transcription and require TC-NER to overcome this.

To investigate the actual rates of NER, we measured illudin S-induced repair DNA synthesis by autoradiography (unscheduled DNA synthesis (UDS)). Even in normal cells we found UDS to be very low: only
Fig. 2. XPC/HR23 complex is dispensable for illudin survival. Survival curves obtained as in Fig. 1 after exposure to illudin S (panel (A) for 72 h and panels (B and D) for 24 h) or to UV (panel C). Panel (A): three different XP-C strains tested against XP-A and normal cells. Panel (B): XP-C survival after a 24 h exposure to illudin. Panel (C): extended UV-exposure protocol mimicking low dose rate; cells were irradiated six times over a period of 72 h, summed up to the total doses indicated. Panel (D): illudin sensitivity of fibroblasts from KO mice. Symbols: ( ) H17040 (normal), ( ) H17009 (XP-A), ( ) H17034 (XP-C), ( ) H17003 (XP-C), ( ) H17010 (XP4LE (XP-C), ( ) DKO (HR23A/HR23B double mutant). Error bars represent S.E. and drawn wherever they exceed the symbol size.

after an exposure to relatively high doses (60 ng/ml for 24 h, about 20 times higher than the D_{10} equivalent) and an extended radioactivity incorporation period during exposure (eight times longer than usually applied with UV), significant UDS above background could be measured in normal fibroblasts. Calculations showed these levels to be at least 30 times lower than what is common for a saturating dose of UV (16 J/m^2), consistent with the estimated low levels of lesions [11]. The UDS we observed clearly represented NER, since it was absent in XP-A cells (Fig. 3E). Despite their defective GG-NER, XP-C cells showed approximately the same level as normal cells, whereas in CS cells, carrying the complementary defect in TC-NER, only marginal levels of UDS were detected, comparable to XP-A cells (Fig. 3E). This result was found in both CS-A and CS-B and strongly contrasts with the characteristic normal UDS levels after UV exposure in these strains (see Fig. 3D and E) [17]. We conclude that the observed (low) UDS activity of normal fibroblasts exclusively represents TC-NER, with no significant contribution of GG-NER.
Fig. 3. Illudin survival depends on transcription-coupled NER. Panels (A and C): sensitivity of CS cells (panel (A)) and UVS (panel (C)) to illudin S. For comparison, dashed lines indicate the UV-sensitivities obtained with these cell strains, on a modified dose scale, such that normal response curves of UV and illudin overlap. Panel (B): recovery of overall transcription after illudin exposure. Cells were prelabelled with $^{14}$C-thymidine, exposed to 0 or 30 ng/ml of illudin S for 3 h, rinsed and pulse-labelled with $^{3}$H-uridine 1, 3 and 21 h later. Relative rates of RNA synthesis are expressed as percentages of treated over mock-treated cells. Symbols in panels (A–C): (□) C5RO (normal), (□) XP25RO (XP-A), (□) XP21RO (XP-C), (□) Kps3 (UVS), (□) CS1BE (CS-A), (▽) CS1AN (CS-B), (□) 94RD103 (UV-resistant trichothiodystrophy). Panels (D and E): unscheduled DNA synthesis induced by illudin S and UV, respectively. Averaged data plotted as a percentage of responses in normal cells and compared to similarly plotted UDS-levels induced by UV.

3.4. Involvement of other global repair pathways

Hydroxymethylacylfulvene (HMAF, Irofulven or MGI-114) is a semisynthetic derivative of illudin S [25], which is currently under clinical trial for tumour therapy [4]. Fig. 4 shows that this agent is about 50 times less toxic to human fibroblasts than illudin S (calculated $D_{10}$ values 50 and 1 ng/ml per 72 h, respectively). However, a 10 times increased sensitivity in XP-A cells, combined with normal XP-C responses were also found with HMAF (Fig. 4), suggesting that this drug acts via the same mechanism as the other illudins.
with known defects in BER and NHEJ pathways (all listed in Table 1). In colony-survival assays, XRCC1-mutant BER-deficient cells as well as three different NHEJ mutants (Ku86, DNA-PKcs, XRCC4) were all as sensitive to illudin as their parental CHO strains and/or mutant strains corrected with a normal copy of their defective gene (data not shown). This resistance is not specific to CHO cell metabolism, since NER-deficient CHO mutants are hypersensitive [11]. We conclude, that the two globally acting repair pathways BER and NHEJ play no significant role in illudin cytotoxicity. These data also suggest that illudin lesions do not cause significant levels of DNA breakage.

3.5. Illudin lesions presumably obstruct replication forks

We have shown earlier, that in cells exposed to illudin S, DNA synthesis is inhibited [2]. To investigate the mechanism of DNA-replication interference we measured the requirement of genes involved in post-replication repair (PRR) pathways. In yeast...
as well as human cells the RAD18/RAD6 protein complex is involved in an early step of PRR [26–28]. Chicken DT40 lymphoma cells with two defective RAD18 alleles are about two times hypersensitive to UV (Yamashita et al., manuscript in preparation). For unknown reasons, chicken lymphoma cells responded to illudin in a highly variable manner, irrespective of the different standardised growth conditions tested. Fig. 5A summarises all our data, indicating that RAD18-deficient cells are hypersensitive to illudin, although experimental variability does not allow a precise estimation of the extent.

Polymerase η (RAD30A) is one of the translesion polymerases active in a downstream branch of the PRR pathway (for overviews see: [29,30]). Absence of this polymerase in XP-variant fibroblasts [31] did not or hardly affect illudin toxicity (see Fig. 1D and 5B) Sensitisation by caffeine, which is a characteristic feature of UV-irradiated pol η mutants [32,33] was also not observed with illudin (Fig. 5B). It follows that the downstream translesion subpathway of PRR is either not required or is accomplished by another translesion polymerase (e.g. pol ζ, pol ξ [34]) which we could not investigate due to lack of mutants.

4. Discussion

Our present detailed data with cell strains from patients having xeroderma pigmentosum and from mice with targeted NER mutations, confirm and extend an early observation in CHO-cells, that NER can repair illudin lesions [11,35]. The overall responses of the non-transformed fibroblasts are highly reproducible over different experiments and cell strains and are found in the same order of magnitude as in relatively resistant tumour cells in culture [9] which appear to rule out large differences in transport and/or metabolic capacities.

Total absence of NER (as occurring in XP-A cells) renders human and mouse fibroblasts about 10 times more sensitive to illudin S, the same enhancement as seen with exposure to UV. This parallelism is retained in other XP complementation groups with NER defects of varying severity. Unexpectedly however, cells defective in components of the HR23/XPC complex, present a notable exception, being normally resistant to the drug. We conclude, that repair factor XPC/HR23 is not used or not required for NER of illudin-induced lesions. The XPC/HR23B protein complex is known to be selectively involved in global-genome repair (GG-NER) [36], where it performs an early function in lesion recognition [37,38] and recruitment of subsequent core NER enzymes [39]. Dispensability of GG-NER was further supported by the independence of XPE, a gene encoding a component of the DDB complex, which plays a complementary role in initiation of GG-NER [40].

According to current models [19,24,41] transcription forks stalled at a lesion in the transcribed strand form a structure that can initiate the so-called transcription-coupled subpathway of NER (TC-NER) without the need of damage recognition by the XPC/HR23B and/or DDB complexes, but still requiring all the other NER factors. We find here that illudin lesions do interfere with transcription, and require TC-NER to recover. Cells defective in specific TC-NER factors (CSA,CSB,UVS) are more sensitive to illudin then they are to UV. The low levels of overall repair (measured as UDS) fully depend on these TC-NER factors. Unfortunately, these low levels and their prolonged induction times have hampered our attempts to perform actual biochemical measurements of transcribed strand-directed repair of actively expressed genes. Since in general, the rate of overall removal of lesions by NER largely depends on the affinity and levels of HR23/XPC [38] and DDB complexes [42], our data are best compatible with the notion that illudin lesions are recognised by GG-NER with very low, undetectable affinity.

While NERable lesions appear to constitute about 90% of potentially lethal damage in fibroblasts, actively transcribed regions represent a much smaller fraction of the genome. It appears that prolonged stalling of transcription forks are the primary cytotoxic entity initiating the apoptotic sequence, as suggested on other grounds earlier [43–45]. Therefore, alternative or additional routes for global repair of illudin lesions could also contribute to protection against illudin. Our data with XRCC1-defective cells indicate that the most common short-patch branch of base excision repair [46,47] is probably not a candidate. The possibility that illudin-lesions lead to significant levels of strand breakage is also considered unlikely, since non-homologous end-joining defects confer no sensitivity to illudin. Taken together, our
Illudin damage not only affects transcription complexes, but inhibits DNA replication as well [2]. We show that there is a strong requirement for RAD18, to overcome such replication blockage. Chicken cells had to be studied here, being the only RAD18-deficient cells currently available. In parallel with TC-NER defective cells (CS-A, CS-B, UVS), they are more sensitive to illudin than to UV. The RAD18/RAD6 protein complex accomplishes an early step in post-replication repair (PRR) [26–28]. A downstream subpathway of PRR involves translesion synthesis by one of a range of recently identified tolerant DNA polymerases [30]. The fully normal responses of XP-variant cells show that polymerase \( \eta \) (with preference for UV CPD-lesions [48,49]) is not the enzyme responsible for this action. However, the involvement of other candidate translesion polymerases (e.g. pol \( \zeta \), pol \( \delta \)) with different specificities [50] was not investigated here yet. A role in illudin cytotoxicity of the alternative RAD5-dependent downstream subpathway of PRR [51] also remains to be checked. Finally, we do not yet know whether there is involvement of other replication-coupled repair modes, such as recombination or mismatch-repair, in recovery from illudin lethality. However, especially with respect to recombinational repair such involvement seems likely, since prolonged stalling of replication forks (e.g. in RAD18-defective cells) is known to increase the frequency of recombination [52]. Moreover, topoisomerase inhibitors, expected to compromise recombinational processing of stalled replication forks, were shown to sensitize cells to illudins [25,53,54].

Information on the nature, chemical structure and stability of illudin-induced DNA lesions, is still lacking. Illudins probably do not produce intrastand crosslinks, in view of moderate sensitivity of ERCC1-deficient cells [11] and data not shown). Our data indicate that global repair pathways appear not to act on the lesion, which is therefore unlikely to cause major helix-distortion. To our knowledge, such an unusual repair pattern has so far only been found with a new class of agents producing adducts exclusively in the minor groove of the DNA helix [55]. The susceptibility to TC-NER was not investigated by these authors, but the exquisite sensitisation in RAD18 mutants which they observed, is fully consistent with our data. Etoxacin 743 offers another case of TC-NER-specificity for minor-groove alkylation [56], although this lesion seems to poison the NER process itself.

In conclusion, we show here that illudins, belonging to a new and promising class of tumour-therapeutic agents, produce a unique type of DNA lesion which is largely ignored by global repair pathways. Only when the lesions become trapped in transcription-complexes or replication-forks, efficient repair mechanisms ensue, such as TC-NER or PRR. As a consequence, we find that gene defects involved in these transcription- and replication-coupled repair pathways confer highly exaggerated sensitivity to the illudins, in comparison to, e.g. UV light. Further studies on more complicated endpoints such as mutagenesis and responses in repair-deficient mice will be needed to expand on this issue.

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