Robust DNA Repair through Collective Rate Control

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Declaration

The applicant, Tim Heinemann, declares that he is the sole author of the submitted dissertation and no other sources or materials from those specifically referred to have been used.

In addition, the applicant declares that he has not applied for permission to enter examination procedure at another institution and this dissertation has not been presented to other faculties and not used in its current or in any other form in another examination.

Heidelberg, November 10,	
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Zusammenfassung

The nucleotide-excision repair pathway removes mutagen-inflicted DNA lesions from the genome. Repair proteins recognize DNA lesions and form multi-protein complexes that catalyze the excision of the lesion and the re-synthesis of the excised part. Imaging the dynamics of fluorescently labeled repair proteins in living human cells has revealed that all factors continuously and rapidly exchange at repair sites. We asked how this dynamic mode of protein-complex assembly shapes the repair process. Measuring repair DNA synthesis in intact cells, we obtained a surprisingly simple result. Over the entire process, the rate is proportional to the amount of DNA lesions, where the proportionality factor is a single 'slow' rate constant. Such kinetic behavior is often regarded as evidence for a rate-limiting step, but we show here that it is an emergent property of the dynamic interplay of many repair proteins. As a consequence, the rate of DNA repair is a systems property that is controlled collectively by the expression levels of all repair factors. Given that transcription in living cells has similar dynamic features - rapidly exchanging components of the transcription machinery and slow bursts of mRNA synthesis - collective rate control might be a general property of chromatin-associated molecular machines.

Summary

DNA repair is indispensable for the intracellular protection against environmental and endogenous damaging agents. This is reflected in an increased susceptibility against cellular aging and cancer development as a consequence to impaired repair. Functional repair is carried out by enzymatic macromolecular complexes that assemble at specific sites on the chromatin fiber. How the rate of these molecular machineries is regulated by their constituent parts is poorly understood. Here we quantify nucleotide-excision DNA repair (NER) in mammalian cells and find that, despite the pathways' molecular complexity, repair effectively obeys slow first-order kinetics. Theoretical analysis indicates that these kinetics are not due to a singular rate-limiting step. Rather, first-order kinetics emerge from the interplay of rapidly and reversibly assembling repair proteins, stochastically distributing DNA lesion repair over a broad time period. Based on this mechanism, the model predicts that the repair proteins collectively control the repair rate. Exploiting natural cell-to-cell variability, we corroborate this prediction for the lesion-recognition factors XPC and XPA. Our findings provide a rationale for the emergence of slow time scales in chromatin-associated processes from fast molecular steps and suggest that collective rate control might be a widespread mode of robust regulation in DNA repair and transcription.

1 Introduction

- 1.1 DNA damage
- 1.1.1 Sources of DNA damage
- 1.1.2 Types of DNA damage
- 1.1.3 Variety of DNA damage repair mechanisms
- 1.2 Experimental research on NER
- 1.2.1 Key components of the NER pathway
- 1.2.2 Expression of fluorescent NER factors

comparison between CPDs and 6-4PP

- 1.3 Mathematical models of NER
- 1.3.1 Metabolic control analysis

2 Mathematical Modelling of Nucleotide Excision Repair (NER)

An essential advantage for the research on Nucleotide Excision Repair (NER) is the ability to inflict UV-induced DNA damages within a locally confined area in the cell nucleus [1]. This technique reformed the investigation of chromatin associated processes whose *in vivo* analysis was, until then, focusing on enzyme mobility and exchange dynamics at steady state performed with photobleaching experiments [2, 3]. In contrast, the enclosed distribution of single stranded DNA damages causes the local activation of a multi-protein repair machinery which in turn allows to study the *de novo* assembly and dissociation kinetics of single repair factors. Exploiting this standard experimental setup we acquired a comprehensive data set comprising the accumulation and dissociation time series of seven individual repair enzymes. In addition, we were able to directly quantify newly synthesized DNA measuring the incorporation of a fluorescently modified nucleotide analog.

Based on this data we developed a mathematical model that is able to explain the connection between the dynamic exchange of individual repair factors at the chromatin on one hand and the overall time scale of the repair process on the other. The exact parametrization of the model was adapted iteratively in correspondence with a profile likelihood estimation leading to identifiable parameters and hence an increased predictive power of the model. As a consequence our model can reliably predict the behavior of an experimentally not accessible DNA intermediate within the repair process.

2.1 Fluorescent time-laps imaging of the NER process

2.1.1 Locally inflicted UV-induced DNA damage

To observe the dynamic behavior of NER factors upon local infliction of DNA damage we applied the experimental setup developed by Moné *et al.* (2001) [1]. Therein, NER competent cells are covered with a polycarbonate UV filter containing pores with a given diameter (cf. Figure 2.1A). For our purposes cells were grown on uncoated 24 mm coverslips before overlaid with a mask including pores with 5 μ m diameter [4]. Figure 2.1B illustrates

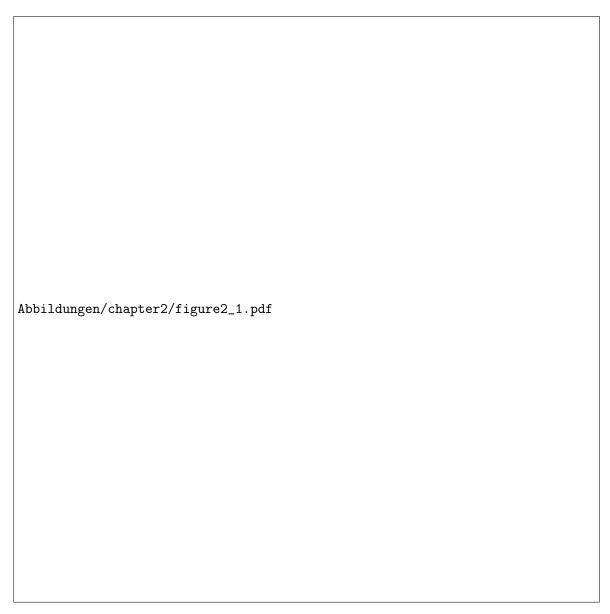


Figure 2.1: Local irradiation leads to spatially confined DNA damage. A) Schematic illustration of the experimental setup. UV-light (purple arrows) is transmitted through a UV filter containing pores with 5 μ m diameter. Local irradiation of the chromatin occurs, if a pore is located above a cell nucleus. (reprint of) B) Distribution of the pore sizes within the UV irradiation mask (n=5008) - ask paul how he did it C) Partially UV-irradiated nuclei of cultivated mammalian cells were made visible with a blue-fluorescent DNA stain. Red fluorescence depicts incorporation of nucleotides due to Nucleotide Excision Repair of damaged DNA.

the realistic deviation of pore diameters due to manufacturing inaccuracies. The actual size distribution has a very small CV of 0.14 indicating a negligible effect on the filter's transmissivity. Filter-covered cells were immediately irradiated with a dose of 100 J/m² of UV-C with a fluency of 3.85 W/m². Due to the specific pore density (4 x 10⁵ pores/cm²) we could observe nuclei containing either one damage spot or no spot at all. Very rarely two spots per cell were present (cf. Figure 2.1C, white arrows indicate damage spot). Consequently, with this technique one can measure NER in damaged nuclei as well as undamaged control nuclei in the same experiment.

2.1.2 Repair factor accumulation and dissociation occur on different time scales

The ability to locally inflict DNA damage with a discrete dose of UV-C allows to study the accumulation and exchange behavior of fluorescently tagged repair proteins under different experimental conditions. In the following we describe the comprehensive dataset acquired by Luijsterburg *et al.* (2010) [5], which represents the basis of our quantitative analysis of the NER process. In total, the kinetics of seven repair factors were measured: i) XPC, the lesion recognition factor; ii) TFIIH, the helicase responsible for DNA unwinding; iii) XPA and RPA, which bind and thereby protect single stranded DNA against cleavage iv) the exonucleases XPF/ERCC1 and XPG performing the incision of the damaged DNA strand and v) PCNA, which loads the DNA polymerase and hence indirectly provides insight into the DNA repair-synthesis kinetic.

Each repair factor is tagged with a green fluorescent protein (GFP) (or its 'enhanced' derivative EGFP) and expressed at physiological levels within the cell nucleus. Before the cells are UV-irradiated at time t = 0 the repair proteins are homogeneously distributed. Immediately afterwards the repair factors accumulate at the damaged chromatin sites, which leads to a higher visible fluorescence intensity (cf. Figure 2.2A). For quantification of the fluorescence intensity image analysis was done by using ImageJ software (NIH Bethesda, MD). Accumulated repair-factor concentrations were determined by multiplying the nuclear reference concentrations (cf. Table 2.1) with the fraction of bound proteins at the damaged DNA:

Bound fraction =
$$(I_{LD} - I_{outspot})A_{LD}/(I_{nucleus} - I_{background})A_{nucleus}$$
 (2.1)

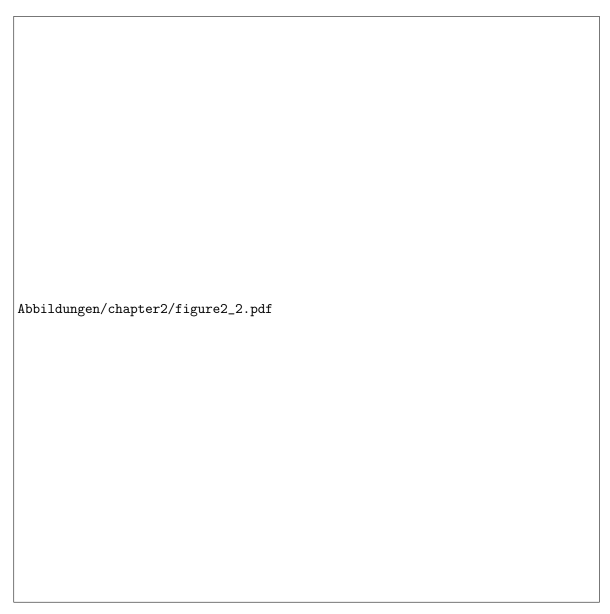


Figure 2.2: Fluorescently labeled NER factors accumulate at locally irradiated nucleus. A) XPC-eGFP accumulation stably expressed in XP-C cells at different time points after local irradiation with UV-C (100 Jm $^{-2}$ through 5- μ m-diameter pores). B) Scheme of a locally UV-irradiated nucleus. Depicted are all regions relevant for signal quantification. C) Time courses of XPC-eGFP (n = 12), XPG-eGFP (n = 5), eGFP-XPA (n = 7), eGFP-PCNA (n = 5) and RPA-eGFP (n = 5) showing their accumulation at the local damage (LD) spot. For consistency only cell nuclei with one single damage spot were used.

where $I_{\rm LD}$, $I_{\rm outspot}$, $I_{\rm nucleus}$ and $I_{\rm background}$ represent the average fluorescence intensities within the locally damaged spot, an equally sized area in the non-damaged nucleus, the whole nucleus and the background, respectively (cf. Figure 2.2B). $A_{\rm LD}$ and $A_{\rm nucleus}$ depict the damaged area and the size of the nucleus. Finally, the concentrations of accumulated protein are calculated assuming a damaged nuclear volume of 0.03 pL.

Protein	Concentration	Bound fraction
XPC	0.140 μ M	13%
TFIIH	0.360 μ M	10%
XPG	0.440 μ M	9%
XPA	1.110 μ M	7%
XPF/ERCC1	0.170 μ M	7%
RPA	1.110 μ M	15%
PCNA	1.110 μ M	20%

Table 2.1: Nuclear concentrations of NER factor in (in \muM) All nuclear quantities are based on published data or on previous estimates [3, 6, 7]. The nuclear concentrations are calculated assuming a nuclear volume of 0.3 pL. The bound fractionis determined by Eqn. 2.1 [5].

During the timespan of DNA repair (cf. Section 1.2) NER factors accumulate towards and then gradually decrease from their plateau levels at different rates (cf. Figure 2.2C). For example the half-life $t_{1/2}$ for XPC- and XPG-EGFP is \sim 1 h whereas \sim 2.5 h for XPA-EGFP [5]. Moreover, PCNA and RPA stay present in the damage spot even after lesion removal. These result show that NER factors engage for hours in the repair process with temporal changes in the molecular composition.

To characterize the interaction between repair proteins and DNA intermediates dwell times were determined in fluorescence loss in photobleaching (FLIP) experiments [5]. Thereby, a large part of the nucleus, away from the local damage spot, is continuously bleached at 100 % laser power (cf. 2.3A, white rectangle). At the same time fluorescent proteins are probed at low laser power elsewhere within the nucleus. Repair proteins dissociating from the local damage spot have a high probability to be bleached before rebinding due to their large diffusivity. Accordingly, for the dwell time of the repair proteins binding instead of diffusion seems to be rate limiting [5].

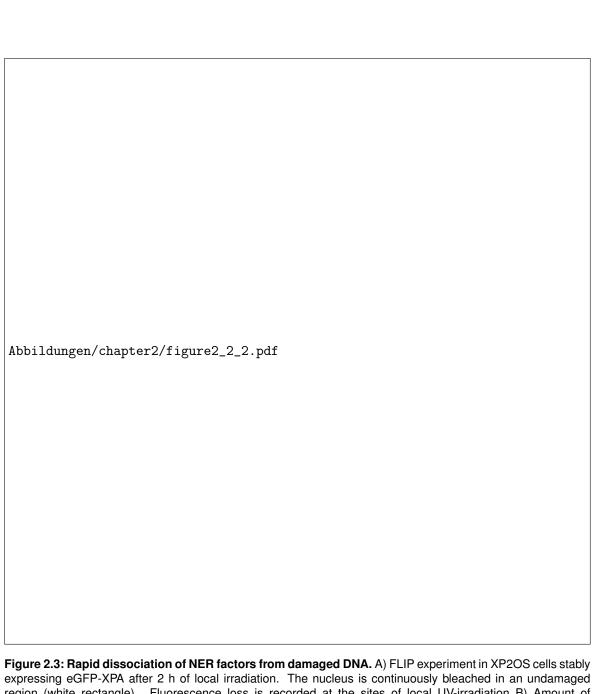
Still, compared to their long overall presence in the range of hours, all NER factors dissociate very quickly from damaged DNA with half-lives of 20 s (RPA), 25 s (XPC), 50 s (TFIIH,

XPG, ERCC1/XPF) and 80 s (RPA) (cf. Figure 2.3B). For PCNA the dissociation is strongly biphasic with half-lives of 10 s and 225 s respectively. To analyze, whether the dwell time of slowly accumulating NER factors changes throughout the repair process resynthesis was stalled by addition of the drugs hydroxyurea (HU) and cytosine- β -arabinofuranoside. While NER factors reaching their plateau level earlier were not affected [5], omitting the repair progression at this late stage mainly slowed down the dissociation of PCNA and RPA (cf. Figure 2.3C). In contrast, XPA's half-life decreased by twofold indicating its higher affinity to repair synthesis intermediates. This shows that the dwell times of NER factors change as repair progresses and suggests that their affinity towards damaged chromatin is defined by the state of the DNA substrate.

2.1.3 Direct measurement of DNA resynthesis

In order to expand our quantitative inspection of the NER pathway in intact mammalian cells we established a protocol for the direct measurement of the repair synthesis process. DNA resynthesis reflects the kinetic of the post-incision repair process, complementing the measure of DNA lesion removal, which captures the systems behavior of the pre-incision steps (cf. section 1.2). To visualize newly incorporated DNA shortly before and after local UV irradiation cells were incubated in microscopy medium supplemented with 10 μ M 5-ethynyl-2'-deoxyuridine (EdU). Due to its excessive presence in solution the DNA polymerase integrates EdU (a thymidine analog) into the new DNA strand instead of the endogenous thymidine (cf. Figure 2.4A). After incubation for the desired time cells were fixed stopping EdU incorporation and subsequently permeabilized. EdU is then tagged with the fluorescent azide (AlexaFluor-555, Life Technologies) forming a covalent bond by click chemistry [8]. Analogous to the quantification of NER factor dynamics EdU intensities were captured with a laser scanning microscope (Zeiss).

Accordingly, incorporated EdU is exclusively present at the locally confined area of damaged DNA, which coincides with the localization of immunostained XPC (cf. Figure 2.4B, upper row). The same result occurs for XP-C cells with stably transfected XPC-eGFP (cf. Figure 2.4B, lower row). Replicating cells could be excluded easily from the analysis due to there prevalent EdU incorporation distributed over the entire nucleus.



2. Mathematical Modelling of Nucleotide Excision Repair (NER)

Figure 2.3: Rapid dissociation of NER factors from damaged DNA. A) FLIP experiment in XP2OS cells stably expressing eGFP-XPA after 2 h of local irradiation. The nucleus is continuously bleached in an undamaged region (white rectangle). Fluorescence loss is recorded at the sites of local UV-irradiation B) Amount of bound NER-factors (XPC-eGFP, TFIIH-eGFP, XPG-eGFP, XPF-GFP, eGFP-XPA, RPA-eGFP and eGFP-PCNA) monitored over time at LD. C) Quantification of FLIP experiments in the absence or presence of HU and AraC for stably expressed eGFP-XPA, RPA-eGFP and eGFP-PCNA

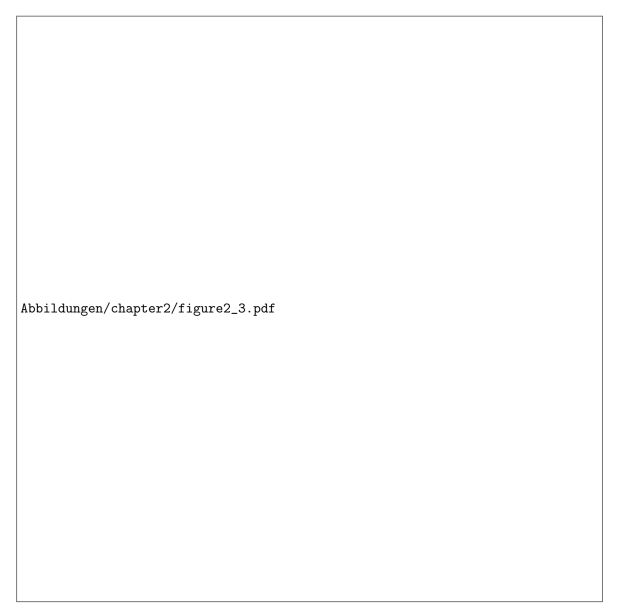


Figure 2.4: EdU is incorporated at sites of local DNA damage. A) Schematic illustration depicting the experimental procedure of EdU incorporation of locally damaged cells. B) Endogenous XPC in Hela cells (upper panel) or stably express XPC-eGFP (lower panel) accumulate in the LD spot and co-localize with incorporated EdU.

2.1.4 Repair rate follows first order rate kinetic

In contrast to the real time measurements for accumulation and dissociation of the NER factors the incorporation of EdU cannot be followed continuously. As mentioned in section 2.1.3 cells have to be fixed and permeabilized before the newly incorporated bases can be fluorescently labeled. Therefore, only the accumulated EdU incorporated in the time interval between UV-irradiation and fixation can be followed. By repeating this procedure for growing time intervals we acquired successively the repair kinetic for newly repaired DNA (cf. Figure 2.5A and B). For each time point we averaged over multiple cells.

We found that EdU incorporation essentially stops after 4 h, which coincides with the removal of 6-4PP (cf. [5] and Figure 2.5B red and blue curve, respectively). This agrees with the observation that NER is not primarily engaged in the removal of cyclobutane pyrimidine dimers (CPD) which are repaired on a much longer timescale. To test whether the availability of EdU is rate limiting we measured the incorporation of EdU in discrete equidistant time intervals after UV-irradiation (cf. Figure 2.5C and Appendix XX). As it turns out the amount per time interval of EdU incorporation is indeed continuously declining and hence the rate of repair synthesis. Moreover the EdU kinetic follows the trajectory of PCNA accumulation as measured by Luijsterburg *et al.* [5] (cf. Figure 2.5D). PCNA in turn is thought to act as processivity factor for the DNA polymerase and remains bound to the DNA [5,9,10]. Taken together, these data establish EdU incorporation as a direct and quantitative measure of DNA repair synthesis in locally damaged cells.

The DNA repair time series characterized by incorporated EdU is fitted by a mono-exponential kinetic (cf. Figure 2.5E):

$$EdU(t) = EdU_{\text{max}}(1 - e^{\lambda t}), \tag{2.2}$$

where EdU(t) and EdU_{max} depict the amount of incorporated EdU and its value at saturation, respectively. The time constant is λ =0.58 (\pm 0.07) h⁻¹. This result indicates that, despite its molecular complexity, 6-4PP removal by NER is a slow first-order reaction with a half-time of 1.2 hours.

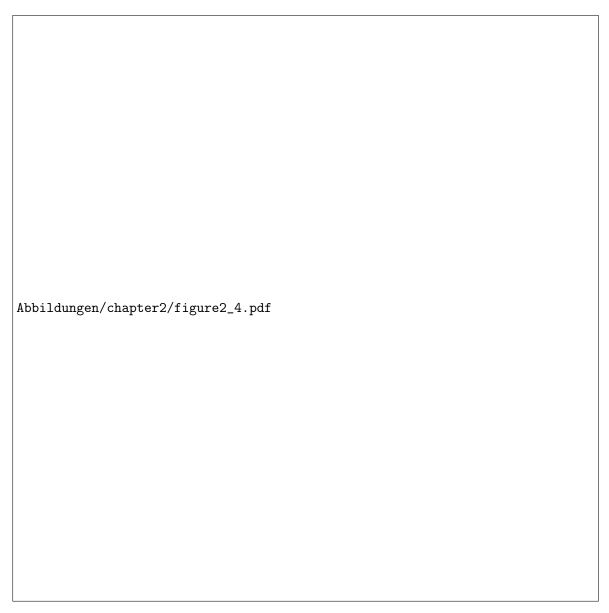


Figure 2.5: EdU incorporation reflects DNA repair synthesis quantitatively. A) EdU signal at various time points after local UC-C irradiation. B) Repair DNA synthesis (EdU, red curve) coincides with damage removal (6-4PP, blue curve) measured previously by Luijsterburg *et al.* [5]. The EdU trajectory represents mean \pm SD (derived from three independent experiments) of DNA repair in n=150 locally damaged cells per time point. C) D) DNA repair synthesis follows PCNA accumulation as measured previously [5]. E) Mono-exponential fit of the EdU kinetic according to Eqn. 2.2.

2.2 Kinetic model of NER

2.2.1 Slow first-order reaction kinetic due to many fast interacting components

To investigate the mechanistic connection between fast NER factor exchange at the DNA template and the overall slow repair time it appears practical to apply an analytical approach. During her Ph.D. thesis Gesa Terstiege together with Thomas Höfer performed this analysis [4,11], considering the complex formation with a simplified model including N repair factors. They examined different scenarios distinguishing random and sequential protein assembly; reversible and irreversible protein binding or mixed mechanisms specific for each DNA repair intermediate. Under appropriate assumptions, the mean repair time of such a process can be generally expressed with:

$$\tau = \underbrace{\frac{1}{k} \sum_{i=0}^{N-1} A_i \left(\frac{l}{k}\right)^i}_{\text{first assembly}} + \underbrace{\frac{1}{\rho} \sum_{i=0}^{N} B_i \left(\frac{l}{k}\right)^i}_{\text{reassembly and reaction}}$$
 (2.3)

where k denotes the pseudo-first order association rate constant of NER factors, l the dissociation rate constant of NER factors and ρ the rate of the repair reaction (cf. Figure **tbd**A, [4]). A_i and B_i differ between random assembly

$$A_i^{\mathsf{rand}} = \sum_{j=1}^{N-i} \frac{1}{i+j} \frac{\binom{N}{j-1}}{\binom{N}{i+j}}, \quad B_i^{\mathsf{rand}} = \binom{N}{i}, \tag{2.4}$$

and sequential assembly

$$A_i^{\rm seq} = N-i, \quad B_i^{\rm seq} = 1. \tag{2.5} \label{eq:2.5}$$

Sequential and random assembly occur on a similar time scale for a number of repair components smaller than 10 considering reversible protein binding (l=k=1) [11], which coincides with the number of core NER factors [5]. The theoretical component number for equal repair times is even higher for increasing reaction rates $(\rho \gg k, l)$. Given that all repair factor dwell times are in the order of \sim 1 min $(k=l=1 \text{ min}^{-1})$ the model was simulated for N = 9 components (cf. Figure **tbm**C). Remarkably, the resulting trajectory followed a single-exponential kinetic with a half time of \sim 1 hour in case of balanced reversibility. In

contrast, for an irreversible process (l/k=0) the repair kinetic resembles a sigmoid time course (cf. Figure**tbm**B). These results suggest that the rapidly exchanging NER factors naturally generate a mono-exponential repair kinetic whereas the slow overall time span can be explained by the stochastic distribution of repair events.

2.2.2 Model structure and parametrization

To examine the relation between rapid repair factor exchange and the slow first-order reaction kinetic we extended this analysis on the basis of a realistic NER model. Conceptually, it follows a model worked out by Luijsterburg *et al.* (2010) [5]. According to this, NER factors bind transiently to DNA repair intermediates thereby forming catalytic complexes that, if complete, perform the next repair step. These are usually irreversible reactions embodying the sequential characteristics of this pathway (cf. Figure **tbm**).

The nature of these distinguishable repair intermediates is widely investigated in biochemical but also in *in vivo* studies [12–15]. Accordingly, the evolved model distinguishes five DNA repair intermediates: i) damaged DNA, unwound DNA, incised DNA, resynthesized DNA and rechromatinized DNA. The molecular state of these intermediates defines the binding affinity for a specific set of repair proteins (cf. Figure 2.7). Table 2.2 summarizes all repair intermediates and lists the corresponding repair factors that show affinity for a particular repair intermediate. Repair factors that catalyze the enzymatic reaction if assembled are indicated. As stated before, FLIP measurements indicate that diffusion is not rate limiting for protein binding and thus not included in the model (cf. Section 2.1.2 and [16, 17]).

A prominent role within the NER pathway belongs to the lesion recognition factor XPC. Theoretical results suggest that the detection of damaged DNA sites by a single element instead of multiple simultaneously is crucial for the efficient initiation of the repair process [18]. Beside this exception the model allows random, non-cooperative binding binding for all NER factors subsequently assembling at the chromatin.

The model structure introduced above and shown in Figure 2.7 was translated into an ordinary differential equation system assuming mass-action kinetics for all protein-DNA interaction processes. Each equation describes the concentration of a single DNA state y_{π}^{R} (cf. Eqn. 2.6) associated with a specific repair intermediate R = I, II,...,V (damaged (I), unwound (II), incised (III), resynthesized (IV), rechromatinized (V)). The index π represents



DNA (V). As indicated, specific tuples of repair proteins bind reversibly to the intermediates. Catalytic reactions,

a binary vector where each position displays the presence or absence of a repair protein p (p ∈ C, T, G, A, F, R, P, where XPC (C), TFIIH (T), XPG (G), XPA (A), ERCC1/XPF (F), RPA (R) and PCNA (P)). $\pi(p)=1$ if protein p is bound and $\pi(p)=0$ if not. The length

of $\pi(p)$ is defined according to the model structure (cf. Table 2.2) with: $\pi_{R=I} = \{\text{C,T}\};$ $\pi_{R=II} = \{\text{C,T,G,A,F,R}\};$ $\pi_{R=II} = \{\text{C,T,G,A,F,R,P}\};$ $\pi_{R=IV} = \{\text{A,R,P}\};$ $\pi_{R=V} = \{\text{R,P}\}.$ The time course of this model is then described by:

$$\frac{d}{dt} y_{\pi}^{R} = \sum_{p \text{ in } R} \eta \left((-1)^{\pi(p)} l_{p}^{R} y_{\pi}^{R} \mid_{\pi(p)=1} + (-1)^{1+\pi(p)} k_{p}^{R} C_{p}(t) y_{\pi}^{R} \mid_{\pi(p)=0} \right) + E(y_{\pi}^{R}), \quad (2.6)$$

where η represents a cooperativity ensuring the exclusion of all cooperative binding events:

$$\eta = \left\{ \begin{array}{rcl} &=& 0 & \text{if} & R = \mathsf{I} \wedge p = \mathsf{C} \wedge \mathsf{T} = 1, \\ & R = \mathsf{I} \wedge p = \mathsf{T} \wedge \mathsf{C} = 0, \\ &=& 1 & \text{else}. \end{array} \right.$$

The kinetics describing protein exchange at the repair intermediates are characterized by the binding constant k_p^R and the dissociation constant l_p^R .

Repair intermediate	Binding proteins	Catalyzed process Required proteins	Remarks	Ref.
(I) Damaged DNA	XPC,TFIIH (3 states)	Unwinding (reaction $lpha$) XPC and TFIIH	Initiation by binding of XPC and subsequent recruitment of TFIIH.	[12] [19] [20] [16] [21]
(II) Unwound DNA	XPC,TFIIH, XPG, XPA, ERCC1/XPF, RPA (64 states)	Dual incision (reaction β) TFIIH,XPG, XPA, RPA and ERCC1/XPF	If the DNA becomes devoid of any protein, it will re-anneal (reaction ε). Dual incision requires the endonucleases XPG and ERCC1/XPF and is stimulated by TFIIH, XPA, RPA and possibly XPC.	[12] [22] [23] [24] [25]
(III) Incised DNA	XPC,TFIIH, XPG, XPA, ERCC1/XPF, RPA, PCNA (128 states)	Repair-synthesis (reaction γ) XPA, RPA and PCNA	PCNA binds to the free 3'-OH group generated by the ERCC1/XPF incision. DNA polymerase is also required (not measured).	[12] [24]
(IV) Resynthesized DNA	XPA, RPA, PCNA (8 states)	Rechromatinization (reaction δ) RPA and PCNA	Accumulation data imply that XPA binds to repaired DNA while the pre-incision proteins do not (Figure 2.2C).	[26] [27] [5]
(V) Rechromatin- ized DNA	RPA, PCNA (4 states)		RPA and PCNA associate with rep. intermediate <i>I</i> , as levels of bound EGFP-PCNA and EGFP-RPA are high up to at least 4h after UV irradiation while other repair proteins are no longer bound.	[19] [5]

Table 2.2: Model assumptions. Adapted from Terstiege et al. (2010) [11]

The free protein concentrations are denoted by $C_p(t)$ representing seven additional differential equations for $p \in \{C,T,G,A,F,R,P\}$:

$$\frac{d}{dt} C_p = \sum_{R=1}^{V} \sum_{\pi} \xi \left(\delta_{\pi(p)1} l_p^R y_{\pi}^R - \delta_{\pi(p)0} k_p^R C_p y_{\pi}^R \right)$$
 (2.7)

Analogous to η , ξ governs the sequential binding of XPC and TFIIH by:

$$\xi = \left\{ \begin{array}{rl} = 0 & \text{if} & R = \mathsf{I} \wedge \mathsf{C} = 0 \wedge \mathsf{T} = 1, \\ & R = \mathsf{I} \wedge p = \mathsf{C} \wedge \mathsf{C} = \mathsf{T} = 1, \\ & R = \mathsf{I} \wedge p = \mathsf{T} \wedge \mathsf{C} = \mathsf{T} = 0, \\ & = 1 & \text{else} \; . \end{array} \right.$$

Finally, if an enzymatic complex has fully assembled at the DNA template it catalyzes the next repair step, which is represented in the model with the term $E(y_\pi^R)$. After damage recognition by XPC damaged DNA is unwound by the helicase TFIIH with the unwinding activity α , whereas XPC acts as a stabilizing/proof reading factor in parallel. Accordingly $E(y_\pi^R)$ translates into the following catalytic reactions for damaged DNA (R=I):

$$E(y_{00}^{\mathrm{I}}) = \ \varepsilon \ y_{000000}^{\mathrm{II}} \quad \text{ and } \quad E(y_{11}^{\mathrm{I}}) = \ -\alpha \ y_{11}^{\mathrm{I}},$$

If all proteins fall off the DNA template due to false damage detection the DNA will re-anneal with the rate ϵ . Otherwise a complex formed by TFIIH, XPG, XPA, XPF, XPA and RPA will eventually promote the excision of the lesion DNA strand leading to the following catalytic reactions for unwound DNA (R = II):

Once the lesion strand is excised with incision rate β the remaining repair steps are irreversible. Incised DNA is resynthesized with the rate γ by the resynthesis complex

XPA-RPA-PCNA. XPA is assumed to assemble at post incision repair intermediates as suggested by experiments with inhibited incision that showed accelerated dissociation FLIP kinetics for XPA [5]. This result is supported by a chromatin immunoprecipitation (ChIP) experiment with antibodies against XRCC1- Lig III showing the co-precipitation of XPA and RPA but not XPC and TFIIH [28]. Evidence for the importance of PCNA and RPA for the resynthesis reaction was shown by Shivji *et al.* (1995) [27]. This results into the following catalytic reactions for incised DNA (R = III):

$$E(y_{1111111}^{\rm III}) \ = \ \beta \ y_{011111}^{\rm II} \quad \text{and} \quad E(y_{0001011}^{\rm III}) = \ -\gamma \ y_{0001011}^{\rm III}.$$

In correspondence to the previously described accumulation measurements (cf. Figure 2.2) only RPA and PCNA stay bound during chromatin remodeling, the last modeled repair intermediate. This leads to the following enzymatic reactions for resynthesized DNA (R = IV) and rechromatinize (R = IV):

$$\begin{array}{lll} E(y_{111}^{\rm IV}) & = & \gamma \; y_{0001011}^{\rm III}, & E(y_{011}^{\rm IV}) = & -\delta \; y_{011}^{\rm IV}, \\ & & & & & & & & & & \\ \mathrm{and} & & & & & & & & & \\ E(y_{11}^{\rm V}) = & & \delta \; y_{011}^{\rm IV}. \end{array}$$

Since all NER factors can bind independently the DNA template for each repair intermediate there are a total of 2^N repair states where N is the number of repair proteins assembling to the particular repair intermediate. Due to the sequential assembly of TFIIH after lesion detection of XPC the number of states for damaged DNA is reduced to $2^2-1=3$ states. For the remaining repair intermediates we derive $2^6=64$ for unwound DNA; $2^7=128$ states for incised DNA; $2^3=8$ states for resynthesized DNA and $2^21=4$ states for rechromatinized DNA. This results in a total of 214 states including seven differential equations for the free NER-factor protein concentrations.

Summing over all repair states and the respective intermediates associated to one repair factor we can simulate the its accumulation kinetic. As initial conditions serve the measured free protein concentrations denoted in Table 2.1 [5, 11] and the initial amount of inflicted damages whose concentration was estimated with $y_{00}^{\rm I}=3.33\mu{\rm M}$. The remaining states

start at zero. To reproduce the FLIP kinetic for a specific repair factor all corresponding dissociation constants were set to zero at the time of maximal accumulation when the FLIP experiment started. Accordingly FLIP kinetics were acquired after 600 s for XPC and ERCCC1/XPF, 900 s for XPG and TFIIH and 2000 s for XPA and 7200 s for PCNA.

2.2.3 Model fitting

To find a realistic parameterization for its complex structure the model can be mapped to m observables represented by the comprehensive data set introduced previously:

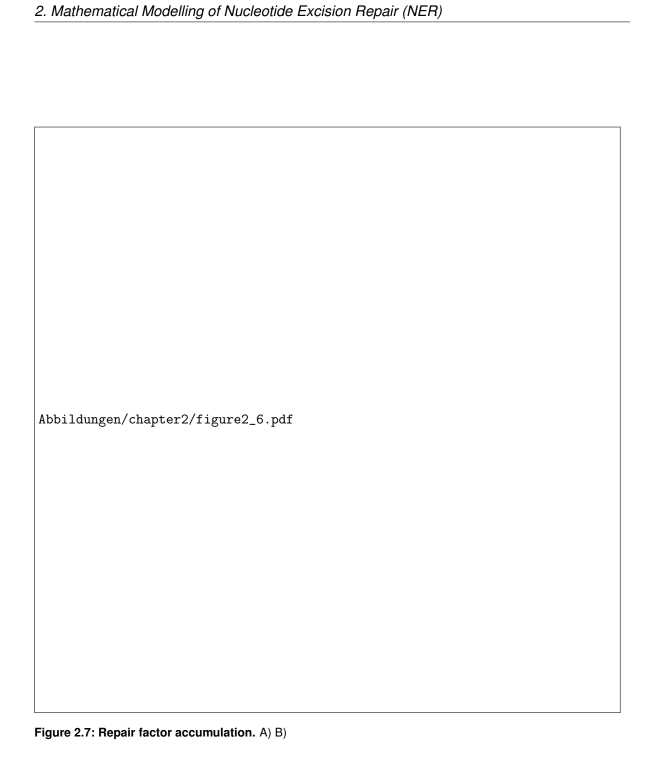
$$y(t_i, \theta) = \tag{2.8}$$

The model structure in turn defines 40 free parameters characterizing binding and dissociation kinetics (20 k_{on} and k_{off} constants, respectively) as well as five catalytic constants determining the sequential repair steps.

For reasons of fitting-speed efficiency we implemented our model into the online-available D2D software environment [29] optimized for MATLAB (2011a, The Mathworks Inc., Natick, MA). The integrated fitting procedure applies a maximum likelihood approach as distance measure, the trust region algorithm LSQNONLIN, which is pre-implemented in MATLAB. The algorithm requires the derivatives of the objective function with respect to the parameters.

These are passed in form of the sensitivity equations:

$$\frac{d}{dt}\frac{dx(t,\theta)}{d\theta} = \frac{\partial f_x}{\partial x}\frac{dx(t,\theta)}{d\theta} + \frac{\partial f_x}{\partial \theta}$$
(2.9)



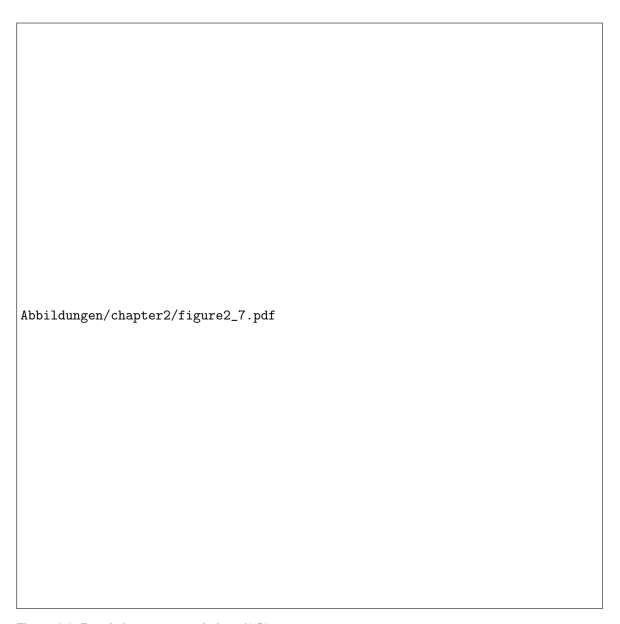


Figure 2.8: Repair factor accumulation. A) B)

- 2.3 Identifiability analysis
- 2.3.1 Prediction Profile Likelihood Estimation (PLE)
- 2.3.2 Refined model structure after PLE analysis

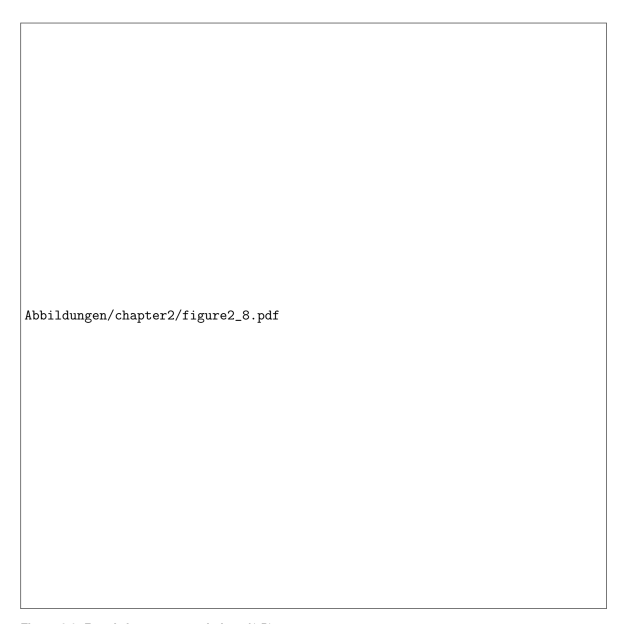
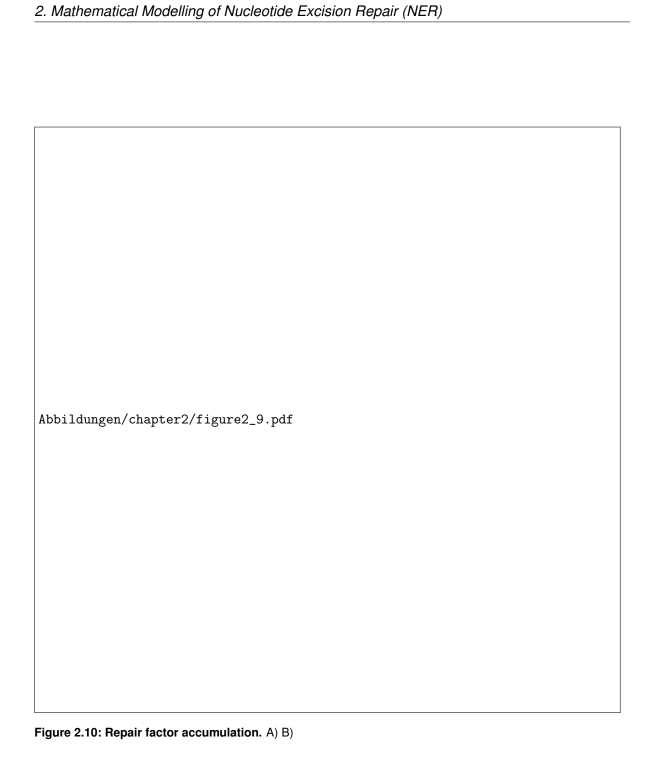


Figure 2.9: Repair factor accumulation. A) B)



3 Control analysis of the DNA repair rate

In the previous chapter we introduced the incorporation of EdU upon UV-irradiation as a quantitative readout for the DNA repair synthesis kinetic (cf. Figure 2.4). We could show that despite the pathway's complexity DNA repair follows a mono-exponential kinetic of first order. Moreover, the repair of single lesions is distributed over a broad time span with 50% of the lesions being repaired after a half-time of \sim 1.2 h. Notably, measurements on the single cell level revealed that also the half-time and hence the rate of repair is highly variable (cf. Figure 2.5B-E). So far the origin of this variability is unknown but also if and how the cell faces this highly fluctuating environment.

In the following chapter we will apply the kinetic NER model and explore the nature of the repair rate variation. Simulating the effect of NER factor variation on the repair rate we find that the repair rate control is distributed among all NER factors. Exploiting the natural variability of expressed repair proteins we can experimentally corroborate the computationally derived finding for the NER factors XPC, TFIIH, XPA, XPF and RPA. Apart from the variability in protein expression the model identifies the initial amount of inflicted DNA damage as major contribution determining the repair rate distribution. Both sources together appear to be sufficient to explain the overall rate variability.

clarify who did what...

3.1 Kinetic NER model predicts collective rate control

3.1.1 Response coefficients

The narrow confidence bounds derived by PLE analysis identified parameters in reasonable biological ranges (cf. Section 2.3.3) and allowed us to use the model for quantitative predictions. On the basis of the simplified model result, indicating that the fast and random enzyme exchange defines the slow first-oder rate kinetics (cf. Section 2.2.1), we wanted to test whether the concept of multi-protein rate control also applies on the realistic NER model. To determine the response of a system to changes in the environment one can calculate the response coefficients \tilde{R}_i [38, 39]. Accordingly, we can quantify the relative change in the

repair rate ν as a function of the relative change in the protein concentration C_i (i = XPC, TFIIH,...).

$$\tilde{R}_i = \frac{\partial \ln \nu}{\partial \ln C_i}.\tag{3.1}$$

The inverse of ν is referred to as characteristic time τ so that Eqn. 3.1 can be rewritten as

$$\tilde{R}_i = \frac{C_i}{\tau^{-1}} \frac{\partial \tau^{-1}}{\partial C_i}.$$
(3.2)

au, in turn, can be directly approximated from the distribution of repaired DNA states $y_{\pi}^{R}(t)$, which include the modeled DNA intermediates for resynthesized and rechromatinized DNA. By taking the ratio between the first $(\mu^{(1)})$ and the zeroth $(\mu^{(0)})$ central moment of the distribution we derive the reaction-specific mean time

$$\tau_R = \frac{\mu(1)}{\mu^{(0)}},\tag{3.3}$$

with

$$\mu^{(m)} = \int t^m y_{\pi}^R(t) dt.$$
 (3.4)

To ensure the convergence of the integral we subtract all repair synthesis states (IV + V) from the initial amount of damages $y_{00}^{I} = 3.33 \,\mu\text{M}$ at t = 0.

$$\tau_{\text{syn}} = \frac{\int (y_{00}^{I}(0) - (\sum_{\pi} y_{\pi}^{IV}(t) + \sum_{\pi} y_{\pi}^{V}(t)) dt) \cdot t dt}{\int y_{00}^{I}(0) - (\sum_{\pi} y_{\pi}^{IV}(t) + \sum_{\pi} y_{\pi}^{V}(t)) dt}$$
(3.5)

Using τ_syn in Equation 3.2 we derive the response coefficients for the repair synthesis rate, which are uniformly small \sim 0.3 and below (cf. Figure 3.1). This result implies that that there is no single repair protein whose effect on the repair speed could be interpreted as rate-limiting. A similar result is obtained for the rate of re-synthesis response coefficients (cf. Appendix), with the characteristic time

$$\tau_{\text{inc}} = \frac{\int_0^\infty t \, \sum_{x=I}^{\text{III}} \sum_{\pi} y_{\pi}^x(t) \, dt}{\int_0^\infty \sum_{x=I}^{\text{III}} \sum_{\pi} y_{\pi}^x(t) \, dt}.$$
 (3.6)

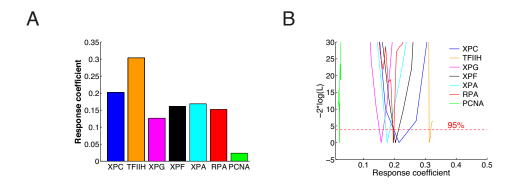


Figure 3.1: Collective control of the repair rate. A) Response coefficients for seven repair factors (XPC, TFIIH, XPG, XPA, XPF, RPA, PCNA) are small and uniformly distributed. B) Prediction profile likelihood indicate small prediction confidence bounds for each response coefficient. 95% threshold is given by the X^2 distribution with one degree of freedom.

Prediction profile likelihood To obtain realistic confidence bounds for the response coefficients we performed a prediction profile likelihood estimation [40, 41]. Thereby the predicted response coefficients $\tilde{R}=F(D_{\text{pred}},\theta)$ are considered as model outcome F for a predicted experimental design $D_{\text{pred}}=(z_{\text{pred}},t_{\text{pred}},u_{\text{pred}})$. D_{pred} specifies a prediction observable z_{pred} at time point t_{pred} given the externally controlled stimulation u_{pred} . As defined in Eqn. 2.8 $z_{\text{pred}}(y(t),\theta)$ comprises a model simulation that can be mapped to experimentally observable quantities. Analogous to Eqn. 2.12 the prediction profile likelihood

$$PPL(\tilde{R}) = \max_{\theta^* \in \{\theta | F(D_{\mathsf{pred}}, \theta) = \tilde{R}\}} L(z^{\dagger} | \theta^*, \tilde{r}) \tag{3.7}$$

is obtained by maximization over the model parameters satisfying the constraint that the model response $F(D,\theta^*)$ after fitting is equal to the considered value \tilde{r} for the prediction \tilde{R} with respect to the measured data z^\dagger . This procedure is repeated for continuous variations of \tilde{R} . The model response can then be expressed as

$$\Delta X_{\theta_l}^2(\tilde{r}) = \min_{\{\theta, \tilde{r} \in \tilde{R}\}} \left(X^2(\theta, \tilde{r}) \right) - \min_{\{\theta\}} \quad \left(X^2(\theta) \right), \tag{3.8}$$

which describes the difference between the global X^2 minimum and the best fit with r included into the objective function. Similar to Eqn. 2.14 we can determine prediction

confidence bounds

$$PCI_{1-a}(D_{\mathsf{pred}}, z^{\dagger}) = \{\tilde{r} | \Delta X_{\theta_l}^2(\tilde{r}) \le Q_{X^2}(1-a, 1)\},$$
 (3.9)

which include the set of predictions $\tilde{R} = F(D_{\text{pred}}, \theta)$ for which -2log(PPL) is below the threshold given by the X^2 -distribution. The PPLs were computed within the d2d-framework [29] using the CVODE package [31] for the numerical integration of the ODEs and the sensitivity equations (cf. Section 2.3.1). By applying the PPL analysis for each response coefficient we find that all of them have small confidence bounds indicating how well the model predictions are determined by the data (cf. Figure 3.1B).

The moderate response predicted by the response coefficients also holds true for larger variations in the repair factor concentration (cf. Figure 3.2). The linear approximation (on which the response coefficients are based) yields a reasonable description for about two-fold concentration decreases or increases (corresponding to a knockdown or overexpression experiment), while for very large decreases the repair rate drops eventually to zero (corresponding essentially to a gene knockout). From this result we can conclude that kinetic NER model predicts a collectively NER factor control of the repair rate. Consequently, the repair pathway appears robust against natural fluctuations in repair protein expression.

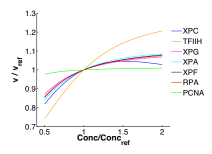


Figure 3.2: Moderate repair rate response against natural NER factor expression variability. Repair rate as a function of concentration changes in individual repair factors.

3.1.2 Exploiting natural variability in protein expression to quantify rate control

To corroborate the model prediction of a distributed repair rate control, we developed an experimental setup for the investigation of single repair factors and their quantitative influence on the repair rate. In particular, we asked whether there is a measurable response in the repair rate due to the natural occurring variability in protein expression. By capturing the integrated nuclear fluorescence intensity of antibody-stained repair proteins we derived the expression values for XPC, TFIIH, XPA, XPF and RPA. (cf. Figure 2.2B). For these five individually measured NER factors the expression variability was quantified by calculating the coefficient of variation (CV; standard deviation divided by the mean), which was on average \sim 0.37 (cf. Figure 3.3A-E). Table 3.1 depicts mean and standard deviation representing at least three biologically independent measurements of the expressed protein cell-two-cell variability.

	XPC	TFIIH	XPA	XPF	RPA
CV:	$\textbf{0.34} \pm \textbf{0.05}$	$\textbf{0.33} \pm \textbf{0.02}$	$\textbf{0.33} \pm \textbf{0.03}$	0.4 ± 0.04	0.44

Table 3.1: Mean and standard deviation of the variability in nuclear XPC, TFIIH, XPA and XPF expression. Distributions for nuclear protein expression were acquired in $n_b(XPC) = 5$, $n_b(TFIIH, XPA, XPF) = 3$ and $n_b(RPA) = 2$ biological independent replicates. Within each measurement between n=250 and n=572 with an average of n=477 cells were analyzed.

To distinguish whether the measured variability is due to differences in nuclear expression or rather superimposed by measurement noise we co-analyzed XPC-eGFP stably expressed in XP-C fibroblasts together with immunofluorescently labeled XPC in the same cell. Both quantities are strong positively correlated (cf. Figure 3.3F) suggesting a large natural variability compared to much lower measurement noise. To quantitatively validate this observation we performed a principal component analysis [42]. Thereby both quantities, XPC-eGFP and antibody-stained XPC intensities, are orthogonally transformed into a new coordinate system were the new transformed variables are linearly uncorrelated. These new variables are referred to as principal components. In a two-dimensional case the variances of both principal components define an error-ellipse as illustrated in Figure 3.3F. From this we estimated a relative measurement error of antibody labeling of ~11% showing that the technique is suitable for quantification of nuclear XPC, TFIIH, XPA, XPF and RPA concentrations.

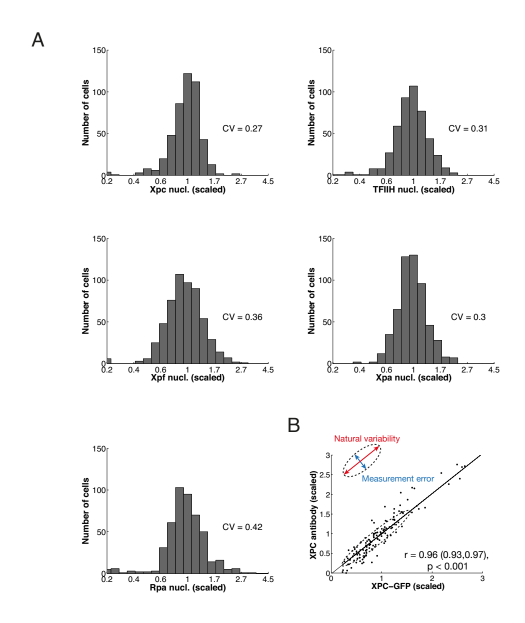


Figure 3.3: Natural variability in nuclear NER factor expression is significantly larger than measurement error A-E) Histograms of nuclear protein concentrations of the antibody stained NER factors XPC, TFIIH, XPF, XPA and RPA (Bildunterschrift: Nuclear und cpt. letters/n nachschauen). Scatter plot of antibody recognized XPC against XPC-eGFP stably expressed in XP-C XPC-eGFP cell lines. The dashed error-ellipse illustrates the proportion of natural variability and measurement noise.

Before exploring the direct relation between protein expression variability and the speed of repair we tested whether higher protein amounts in the nucleus correlate with the accumulation of NER factors in the locally damaged area. In correspondence to previous findings for XPG [5] there is a significant positive correlation between the nuclear XPC-eGFP expression and its local accumulation at the DNA lesions (cf. Figure 3.4A) thirty minutes after UV irradiation. The same holds true for XPA [4], TFIIH, XPF and RPA in narrow confidence bounds (cf. Appendix **tbm**). Consequently, we conclude that the DNA lesions are not saturated so that a higher NER factor concentration could potentially accelerate the repair rate.

In fact, the nuclear protein concentrations for all five antibody-stained repair factors and the amount of incorporated EdU after one hour were significantly correlated (cf. Figure 3.4B-F). Remarkably, the dependency between both quantities, characterized by the slop of the regression line is in the same order of magnitude as predicted computationally by the response coefficients (cf. Figure 3.1A). Also among each other, correlation and regression slope fell into the same range. This agrees with the *in silico* finding that the kinetic control of the measured repair factors is distributed among them and that the rate of repair synthesis is robust against natural variations in the repair protein expression.

Correlation analysis The correlation analysis was performed in MATLAB (2012a, The Mathworks Inc., Natick, MA) using the inbuilt corrcoef function, which also returns p-values indicating the significance of the correlation coefficient. The p-values are determined by a t-test, where the null-hypothesis states that the true correlation between two variables equals zero and that for its observed value r the quantity

$$t = \frac{r}{\sqrt{\frac{(1-r^2)}{N-2}}}\tag{3.10}$$

follows approximately a t-distribution for N-2 degrees of freedom and a sample size N [43,44]. A low p-value suggests the rejection of the null-hypothesis and thus, demonstrates the significance of the correlation.

confidence bounds were derived by bootstraping ...

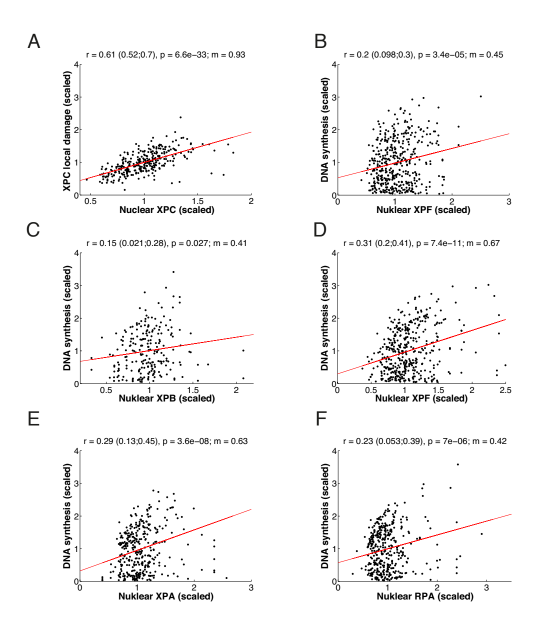
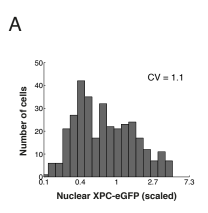


Figure 3.4: Blubb. A) B)

3.1.3 NER Variability comparable between complemented patient cell lines and native repair



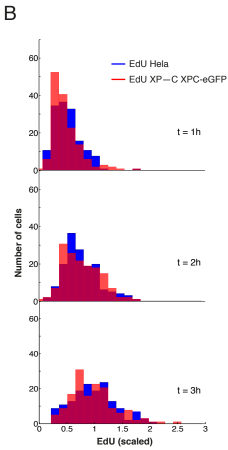


Figure 3.5: Blubb. A) B)

3.2 Sources for repair rate variability

3.2.1 Variable NER factor expression and inflicted lesions account for the distribution of repair rates

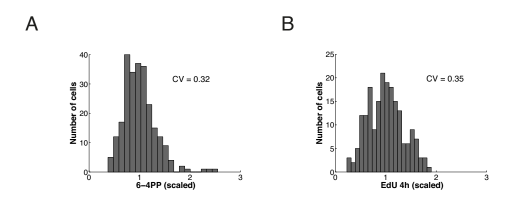


Figure 3.6: Blubb. A) B)

3.2.2 Imbalanced contribution

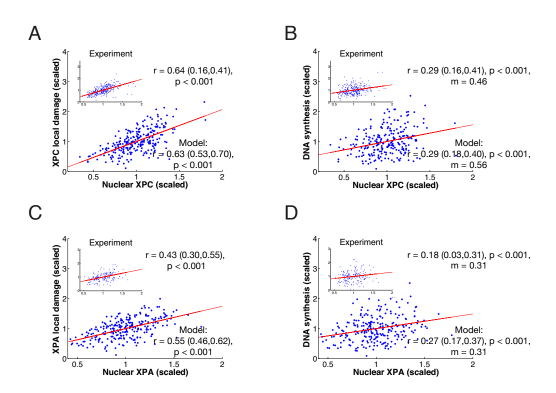


Figure 3.7: Blubb. A) B)

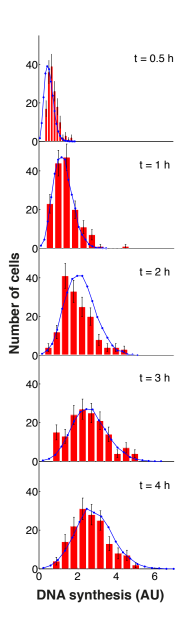


Figure 3.8: Blubb. A) B)

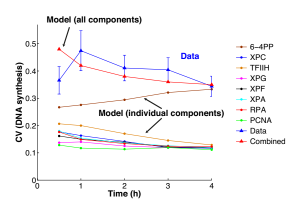


Figure 3.9: Blubb. A) B)

4 Co-expression of NER repair factors

The previously described model-aided analysis of the DNA repair process revealed the link between the emergent phenomenon of rapidly exchanging and transiently interacting NER components with the experimentally observed slow first-order kinetics of repair. An important functional consequence of this kinetic design is that the control of the repair rate is shared by all repair factors. This manifests in the mathematical prediction of uniformly distributed response coefficients, which quantify the relative change of the repair rate in answer to changes in the nuclear repair protein concentration. Exploiting the natural variability in NER factor expression, their moderate control on the rate of repair could be experimentally resolved for different repair phases. These findings, however, were made under the assumption of a functional independence of the individual NER components. Effects resulting from co-regulated NER factor expression or protein degradation were so far neglected as a first approximation.

To test this assumption, in the following chapter we will investigate a potential cross-correlation among the involved repair factors experimentally. Surprisingly, we find that the nuclear expression of five pairwise measured repair factors (XPC, TFIIH, XPA, XPF and RPA) is indeed strong positively correlated, whereas there is no correlation with the repair-independent cell cycle marker **Ki67**. This result suggests an additional control mechanism orchestrating NER factor expression on the transcriptional or translational level. - sentence about who did what

4.1 Co-staining experiments

Using fluorescence microscopy for the quantitative analysis of the NER process proved to result in accurate measurements of the nuclear expression and UV-induced repair dynamics of the individual repair factors at a high signal to noise ratio. This applies for the stably transfected fluorescently tagged repair protein expression as well as for an immunocytochemistry approach with an indirect antibody-labeling of the measured NER factors. The latter technique has the advantage that once a specifically binding antibody is established it can be applied on all cell types containing the desired antigen. On

this basis, indirect labeling of the primary antibody with a secondary antibody allows for the combinatorial tagging of multiple antigens restricted only by cross-reactions between antibodies originating from the same host species [45, 46]. Following such a protocol we established five single cell double stainings (cf. Table 4.1 using the primary and secondary antibodies listed in Table **tbm**.

	XPC	TFIIH	XPA	XPF	RPA
XPC	Χ	Χ	Χ	Χ	Χ
TFIIH		-	_	Χ	-
XPA			-	Χ	Х
XPF				-	-
RPA					_

Table 4.1: Mean and standard deviation of the variability in nuclear XPC, TFIIH, XPA and XPF expression.

Additional two cross-correlation measurements were possible because of a directly labeled monoclonal mouse XPC-antibody. After an initial indirect labeling step for one of the other four repair factors no further secondary antibody is needed. Hence, there is no cross-recognition for primary antibodies from the same species. Moreover, the direct labeling permits a double staining for XPC with two individual fluorophors, which offers a precise estimate of the measurement error (cf. Section 3.1.2).

The cross-correlation analysis was performed in human diploid female fibroblasts which were grown to confluency on coverslips. Analogous to the description in Section 2.1.1 cells were UV-irradiated locally with a dose of 100 J/m². Pre-incubated with serum-free medium containing 10 µM EdU cells were allowed to repair for 60 minutes in an incubator. After the subsequent direct or indirect antibody labeling of two selected repair factors cells were additionally incubated with a DAPI solution, which visualizes the cell's chromatin and thus the contour of the nucleus. For each double staining microscopic 3-dimensional imaging was conducted on a Leica TCS SP5 II confocal microscope. All images were analyzed following the protocol presented in Section 2.1.2. Segmentation of the nucleus was performed on the DAPI signal, due to the large signal-to-nose ratio in this channel (cf. Appendix **Appendix**

and Figure 4.1A). The segmented region was then used to quantify the signal emitted by the secondary antibodies at 488 and 647 nm (cf. Figure 4.1B and C). For the same reason segmentation of the locally UV-irradiated chromatin was done with the EdU signal and then projected onto the signal of accumulated protein (cf. Figure 4.1D).

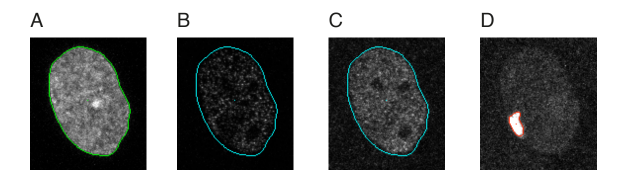


Figure 4.1: Blubb. A) B)

4.1.1 Nuclear expression of NER factors is strongly correlated

To test whether the antibody co-staining experiment is suitable for the cross-correlation analysis we measured XPC with a directly labeled antibody and an indirect labeling and correlated both signals (cf. Figure 4.2). Applying PCA as described in Section 3.1.2 the measurements error was determined with ... This lies in the same range as the combined approach, with one protein stably expressed and the other indirectly labeled. Hence, the method

4.1.2 Flow cytometry verification

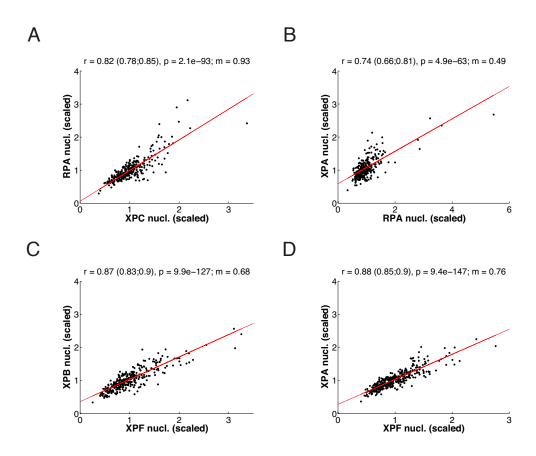


Figure 4.2: Blubb. A) B)

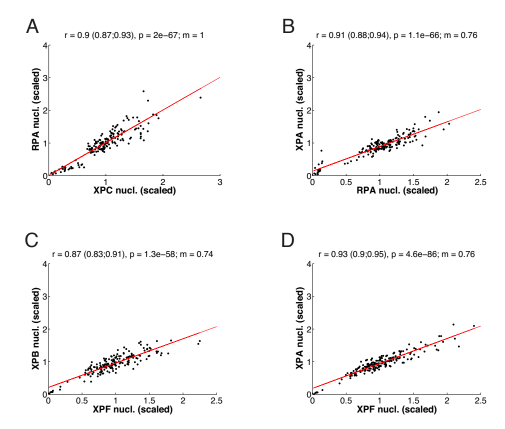


Figure 4.3: Blubb. A) B)

5 Discussion

- discuss difficulties during antibody staining

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7 Appendix

		dissociation	

Value	XPC	TFIIH	XPG	XPF	XPA	RPA	PCNA
Concentration (µM)	0.140	0.360	0.440	1.110	0.170	1.110	1.110
Damaged DNA							
k _{on} (μM ⁻¹ s ⁻¹)	0.025	0.2	NA	NA	NA	NA	NA
	(0.013;0.036)	(0.152;0.288)					
k _{off} (s ⁻¹)	0.231 (0.136;0.36)	0.01 (0.009;0.012)	NA	NA	NA	NA	NA
K _d (μM)	9.35	0.052	NA	NA	NA	NA	NA
ι (α (μινι)	(3.46;16.09)	(0.044;0.072)	IVA	IVA	IVA	INA	IVA
Unwound DNA	(0,,	(0.0,0.0)					
k _{on} (μM ⁻¹ s ⁻¹)	0.022	0.049	0.037	0.0056	0.116	0.018	NA
ιτοη (μινί · 3 ·)	(0.016;0.025)	(0.04;0.06)	(0.036;0.039)	(0.0054;0.0059)	(0.109;0.125)	(0.016;0.02)	IVA
$k_{off}(s^{-1})$	0.019	0.01	0.0146	0.0137	0.017	0.022	NA
οπ (Ο)	(0.019;0.019)	(0.009;0.012)	(0.0140;0.0149)	(0.0134; 0.0141)	(0.0167;0.0172)	(0.0213;0.0224)	
$K_d(\mu M)$	0.864	0.204	0.395	2.446	0.147	1.222	NA
· · · · · · · · · · · · · · · · · · ·	(0.635;1.006)	(0.163;0.259)	(0.373;0.419)	(2.344; 2.596)	(0.138;0.158)	(1.048;1.36)	
Incised DNA	(,	(,-	(/ /	(- ,,	(,,	(,,	
$k_{on}(\mu M^{-1} s^{-1})$	0.022	0.049	0.037	0.0056	0.116	0.018	0.008
·········· • /	(0.016;0.025)	(0.04;0.06)	(0.036;0.039)	(0.0054;0.0059)	(0.109;0.125)	(0.016;0.02)	(0.0066;0.0111
$k_{off}(s^{-1})$	0.019	0.01	0.0146	0.0137	0.017	0.022	0.0031
Oli (-)	(0.019; 0.019)	(0.009; 0.012)	(0.0140; 0.0149)	(0.0134; 0.0141)	(0.0167; 0.0172)	(0.0213; 0.0224)	(0.003; 0.0032)
$K_d(\mu M)$	0.864	0.204	0.395	2.446	0.147	1.222	0.388
44 /	(0.635;1.006)	(0.163; 0.259)	(0.373; 0.419)	(2.344; 2.596)	(0.138; 0.158)	(1.048;1.36)	(0.319;0.538)
Resynthesized DNA	, , , ,	, , ,	, , ,	, , ,	, , ,	, , ,	, ,
$k_{on}(\mu M^{-1} s^{-1})$	NA	NA	NA	NA	0.022	0.032	0.004
On (P					(0.021;0.025)	(0.025; 0.037)	(0.0038;0.005)
$k_{off}(s^{-1})$	NA	NA	NA	NA	0.0052	0.035	0.0026
011()					(0.0051;0.0053)	(0.031;0.042)	(0.0025;0.0027
$K_d(\mu M)$	NA	NA	NA	NA	0.236	1.167	0.605
a (r·)					(0.222;0.27)	(0.924;1.521)	(0.531;0.747)
Rechromatinized DN	NA A				, , ,	, ,	,
$k_{on}(\mu M^{-1} s^{-1})$	NA	NA	NA	NA	NA	0.065	0.396
- /						(0.056;0.075)	(0.359;0.457)
k _{off} (s ⁻¹)	NA	NA	NA	NA	NA	0.035	0.061
··· /						(0.031;0.04)	(0.056; 0.067)
$K_d(\mu M)$	NA	NA	NA	NA	NA	0.538	0.154
/						(0.438; 0.645)	(0.134; 0.182)

Table SII. Values of the enzymatic rate constants

Enzymatic rate		k _{cat}
		s ⁻¹
Unwinding α	19.9	(>0.2)
Resynthesis γ	25.5	(>1.5)
Rechromatinization δ	0.001	(0.001;0.0011)
Reannealing ϵ	5.3	(>0.9)