Human base excision repair complex is physically associated to DNA replication and cell cycle regulatory proteins

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ABSTRACT

It has been hypothesized that a replication-associated repair pathway operates on base damage and single strand breaks (SSB) at replication forks. In this study, we present the isolation from the nuclei of human cycling cells of a multiprotein complex containing most of the essential components of base excision repair (BER)/SSBR, including APE1, UNG2, XRCC1 and POLβ, DNA PK, replicative POLα, δ and ε, DNA ligase 1 and cell cycle regulatory protein cyclin A. Co-immunoprecipitation revealed that in this complex DNA repair proteins are physically associated to cyclin A and to DNA replication proteins including MCM7. This complex is endowed with DNA polymerase and protein kinase activity and is able to perform BER of uracil and AP sites. This finding suggests that a pre-assembled DNA repair machinery is constitutively active in cycling cells and is ready to be recruited at base damage and breaks occurring at replication forks.

INTRODUCTION

Both DNA replication and repair are performed by multiprotein assemblies and share common features. DNA repair must be coordinated with DNA replication in order to avoid fixation of DNA damage into heritable mutations. A fundamental level of cross-talk between DNA replication and DNA repair is ensured by the fact that the enzymes active in replicating DNA during S phase are also needed to synthesize new stretches of DNA during various types of repair including nucleotide excision repair (NER), mismatch repair (MMR), single-strand DNA break repair (SSR) and base excision repair (BER). An additional level of coordination is needed to achieve integration of the DNA repair and DNA replication protein networks within the highly sophisticated cell cycle regulatory machinery. Indeed, many components of the DNA replication machinery associate with other factors such as cyclins/Cdkks in dynamic multiprotein complexes that regulate cell cycle progression. The so-called cyclin-dependent protein kinase (Cdk)-driven ‘replication switch’ model predicts that cyclin/Cdk complexes function both to activate initiation complexes assembled at the origins and to inhibit further complex assembly during S-phase, thus preventing unscheduled re-replication (reviewed in (1)).

BER counteracts the cytotoxic and mutagenic effects of most endogenously produced DNA damage. Its role must be critical when this type of damage is produced or persists at replication forks. In BER, specific DNA glycosylases are responsible for base removal followed by formation of a single strand break (SSB) by an AP endonuclease. SSB that arise directly from sugar damage usually possess non-conventional termini that need further processing to allow completion of SSB repair (SSBR). In both BER and SSBR the resulting gap is filled by DNA polymerase (POLβ (short-patch repair) or by POLβ/δ/ε (long-patch repair). Additional players in long-patch repair are replication factor (RF) C, proliferating cell nuclear antigen (PCNA) and flap endonuclease-1 (FEN1). The final ligation step is then operated by the XRCC1/DNA ligase IIIα (LIG3) complex or DNA ligase I (LIG1) in the short- and long-patch pathways, respectively (reviewed in (2)). Interestingly, PCNA, that is involved in the dynamic assembly and disassembly of the DNA replication machinery, has been shown to interact with several BER/SSBR proteins such as adenine DNA glycosylase (MYH) (3), uracil DNA glycosylase UNG2, 5′ AP endonuclease APE1, XRCC1, POLβ, POLδ, FEN1and LIG1 (reviewed in (4)). Based on these findings, it has been hypothesized that there is a BER/SSBR
pathway that is coordinated with DNA replication and employs proteins like PCNA, FEN1, LIG1 and POLδ/ε that are in common with the replication machinery (5–7).

A growing body of evidence indicates that BER/SSBR proteins are regulated by post-translational modification and make physical interactions with components of other DNA transaction pathways (reviewed in (8)). One of the most compelling evidence of BER regulation via post-translation modifications is the phosphorylation in vitro and in vivo by Casein Kinase 2 of one of the central players of BER, the scaffold protein XRCC1 (9). This phosphorylated form promotes a more efficient SSBR. Another example is the homeostatic regulation of BER by a p53-activated form, UNG2 (10). This phosphorylation is endowed with a protein kinase activity and is able to employ proteins like PCNA, FEN1, LIG1 and POLδ/ε that are in common with the replication machinery (5–7).

It should be taken into account that most of our knowledge about BER has been derived from studies carried out in vitro by using mammalian cell extracts or purified proteins and synthetic DNA molecules containing single lesions. However, an open question is how cross-talk between DNA replication and DNA repair machineries is achieved at the cellular and molecular level. The analysis of the protein–protein interactions within BER proteins and between BER and other pathways occurring in the cell is a prerequisite to better understand the regulation of the DNA repair processes in the context of the cell cycle.

In this study, we present the isolation from the nuclei of human cycling cells of a complex containing most of the essential components of BER physically associated to cyclin A and to DNA replication proteins. This complex is endowed with a protein kinase activity and is able to perform BER of uracil residues as well as of apurinic/apyrimidinic (AP) sites via both short- and long-patch BER. The fact that this complex was isolated from human cells in the absence of any DNA-damaging treatment, suggests that a preassembled BER machinery is constitutively active in the cycling cells and is ready to be recruited to the site of damage likely to occur at the replication forks.

**MATERIALS AND METHODS**

**Chemicals**

Restriction enzymes and T4 polynucleotide kinase were from New England Biolabs. T4 DNA polymerase holoenzyme, T4 single-stranded DNA binding protein, T4 DNA ligase and dNTPs were purchased from Roche Molecular Biochemicals. Both oligodeoxyribonucleotides containing a single uracil: 5′-GATCCTCTAGAGUCGA CCTGCA-3′ (for preparation of circular duplex DNA substrate) and 5′-GATCCTCTAGAGUCGACCTGCAG GCATGCA-3′ (for incision assay), were synthesized by MWG-Biotech AG. Ugi was kindly provided by S.E. Bennett (Oregon State University, Corvallis, OR, USA). [γ-32P] ATP was from GE Healthcare and [α-32P] dCTP and [α-32P]dTTP were from Perkin Elmer.

Nitrocellulose membranes (HybondECL) were from GE Healthcare. GF/C filters were from Schleicher & Schuell. Olomucine and aphidicolin were from Sigma. All other reagents were purchased from BioRad, Sigma, Fluka, Gibco and BDH.

**Cells and media**

HeLa cells were grown in DMEM supplemented with 10% FCS, 50 μg/ml gentamycin and 2 mM l-Glutamine, at 37°C and 5% CO2. Cells were pelleted by centrifugation and stored in aliquots at −80°C until used.

SV40 transformed wild-type and POLβ-null mouse embryonic fibroblasts (MEFs) (a gift from Dr S.H. Wilson, NIEHS, Research Triangle Park, NC) were cultured in DMEM supplemented with Glutamax-1, 10% FCS, 100 U/ml penicillin, 100 μg/ml streptomycin and 80 μg/ml hygromycin at 34°C and 10% CO2.

**Protein purification**

The purification was performed essentially as described (11), with the following modifications. After the size exclusion chromatography, the pooled fractions were loaded onto a hydroxyapatite column, equilibrated in Buffer F (25 mM Bis-Tris pH 6.6, 50 mM NaCl, 0.5 mM ATP, 1 mM DTT, 5% glycerol, 1 mM PMSF and protease inhibitors). The column was eluted with a linear gradient from 0.1 to 1 M phosphate buffer pH 6.6. The active fractions were pooled, diluted in Buffer F to bring the potassium phosphate below 0.1 M and then loaded onto a Mono S column (Pharmacia) equilibrated in Buffer F. The column was eluted with a linear gradient from 0.1 to 1 M NaCl.

**Immunoprecipitation experiments**

The Mono S fraction 15 (5 μg) was adjusted to 10 mM Tris-HCl pH 7.4, 50 mM NaCl, 2.5 mM MgCl2 and protease inhibitors and then incubated for 1 h at 4°C with protein A-agarose beads (BioRad) equilibrated in the same buffer and previously coupled to the appropriate antibodies, as indicated in the figure legends. After centrifugation the pellet was washed with equilibration buffer containing 80 mM NaCl and the immunoprecipitated material was analysed by western blot.

Wild-type or POLβ-null MEFs (4 × 107) were lysed in the presence of 10 mM Tris-HCl pH 7.4, 150 mM NaCl, 2.5 mM MgCl2, and protease inhibitors for 30 min on ice and then homogenized by dounce. After centrifugation at 20,000×g at 4°C, the supernatant (crude extract) was diluted two times in the same buffer without NaCl and incubated for 1 h at 4°C with protein A-agarose beads (BioRad) equilibrated in the same buffer and previously coupled to the appropriate antibodies, as indicated in the figure legends. After centrifugation the pellet was washed with the same buffer containing 80 mM NaCl and the immunoprecipitated material was analysed by western blot.

**Native gel electrophoresis**

Mono S fraction 15 (5–10 μg) was resolved on a 5% native polyacrylamide gel in the absence of SDS and
β-mercaptoethanol in sample loading and running buffers, at 4°C. Samples were transferred to nitrocellulose membrane at 12 V for 12–16 h at 4°C and subjected to western blot analysis.

**Gel filtration**

Mono S fraction 15 (20 μg) was applied to a Superdex200 10/300 GL gel filtration column equilibrated with 10 mM Tris-HCl pH 7.4, 80 mM NaCl, 2.5 mM MgCl₂ and protease inhibitors. The column was eluted with the same buffer and the eluted fraction analysed by dot blot immunoassay and western blot. Molecular weight markers used for column calibration were ferritin (440 kDa), catalase (232 kDa), lactate dehydrogenase (140 kDa) and BSA (66 kDa).

**Protein kinase assays**

The assay was carried out in final volume of 10 μl containing 0.15 μg of Mono S fraction 14, 50 mM Tris-HCl pH 7.5, 10 mM MgCl₂, 0.5 μM [γ-32P] ATP (3000 Ci/mmole). Reactions were supplemented, when indicated, with 0.4 mg/ml histone H1, 0.5 mM Olomucine or recombinant p21cip/WAF. Samples were incubated for 20 min at 37°C and loaded onto a 10% SDS-PAGE. Gels were quantified by PhosphorImager (Typhoon, GE Healthcare).

**Antibodies**

Anti-FEN1, anti-UDG, anti-POLα, POLδ, POLε, anti-cyclin B and E, anti-MCM7 and anti-APE1 were from Santa-Cruz Biotechnology. Anti-cyclin A and anti-phosphoSer/Thr were from Sigma. Anti-XRCC1 was from Trevigen. Anti-DNA PK was from Calbiochem. Anti-POLβ was kindly provided by S.H. Wilson (NIEHS, Research Triangle Park, NC, USA) and anti-POLβ by T.A. Kunkel (NIEHS, Research Triangle Park, NC, USA). Anti-LIG1 was a kind gift of A. Montecucco (IGM-CNR, Pavia, Italy) and anti-POLα was kindly provided by U. Hübscher (University of Zürich-Irchel, Switzerland).

**Preparation and characterization of circular duplex DNA substrates**

Closed circular DNA molecules containing a single lesion were produced as described previously (12) by priming single-stranded (+) pGem-3Zf(+) DNA (Promega) with the oligonucleotides containing the lesion of interest. In vitro DNA synthesis was performed by using T4 DNA polymerase holoenzyme, single-stranded DNA binding protein, dNTPs and T4 DNA ligase. Closed circular DNA duplex molecules were purified by cesium chloride equilibrium centrifugation. Plasmid DNA molecules containing a single uracil residue were digested with E. coli uracil DNA glycosylase (Trevigen) to produce abasic sites.

**Repair assay**

Repair reactions were carried out as described in (12). Briefly, reaction mixtures (50 μl) contained 40 mM HEPES/KOH (pH 7.9), 75 mM KCl, 5 mM MgCl₂, 0.5 mM dithiothreitol, 50 μM of each dNTP, 2 μCi of [α-32P] P dCTP or [α-32P] P dTTP as indicated, 2 mM ATP, 40 mM phosphocreatine, 2.5 μg of creatine phosphokinase (type I, Sigma), 3.4% glycerol, 18 μg of bovine serum albumin and 5 μl of the isolated multiprotein complex (Mono S fraction 14, 1 μg of total proteins) were incubated 1 h at 30°C. Depending on the labelled dNTP used in the repair mix, the concentration of the corresponding cold dNTP was decreased to 5 μM. When the assay was performed in the presence of aphidicolin, a concentration of 300 μM was used. The repair products were digested with appropriate restriction enzymes, resolved on a denaturing 20% PAGE and quantified by electronic autoradiography (Instant Imager, Packard).

**Incision assay**

(γ-32P)-labelled 30-mer oligonucleotides containing a single uracil were annealed with the appropriate complementary oligonucleotide to obtain U/A-containing duplexes. Following 1 h incubation at 37°C with the complex (Mono S fraction 14, 1 μg of total proteins) in 25 mM Tris-HCl pH 7.6, 1 mM EDTA, 50 mM NaCl, the incision products were incubated for 10 min at 95°C in the loading buffer (formamide 95%, EDTA 20 mM, bromophenol blue 0.05%, xylene cyanol FF 0.05%), resolved on a denaturing 20% PAGE and quantified by electronic autoradiography (Instant Imager, Packard).

**RESULTS**

**Isolation of a multiprotein complex from HeLa cells containing DNA replication proteins, cell cycle regulatory factors and base excision repair proteins**

HeLa cell nuclei were isolated and fractionated according to a procedure already established in our laboratory to isolate multiprotein complexes (11). During the fractionation procedure, DNA polymerase activity was measured using three different assays, one specific for POLα (activated DNA) another one for POLδ and POLε (poly(dA)/oligo(dT) in the presence of PCNA) and a third one for POLα and POLδ (poly(dA)/oligo(dT) in the absence of PCNA). Co-purification of other DNA replication and cell cycle regulatory factors was monitored by a dot blot immunoassay. Figure 1 shows the elution profile of the last purification step (Mono S column). A peak of DNA polymerase activity was observed between fraction 10 and 20 with both assays (Figure 1A). Further characterization revealed the presence of 3′–5′ exonuclease activity, DNA helicase activity and histone H1 phosphorylation activity (data not shown and Figure 2). The presence of POLα and POLδ and, in addition, LIG1 and cyclin A in the same fractions was confirmed by dot blot (Figure 1B). A slightly different elution profile was revealed at the tails of the peak (fractions 10–12 and 20–24, Figure 1) between cyclin A and the other proteins. This could be either due to different isoelectric points and/or different sensitivities of the antibodies used. The central peak fractions (14 and 15) were used for all the subsequent experiments. Next, the isolated complex (Mono S fraction 14) was analysed with a panel of poly- and monoclonal antibodies directed against a number of DNA replication proteins.
and repair proteins. As shown in Figure 3, western blot analysis revealed the presence in the same fraction of four different DNA polymerases (POLα, POLβ, POLδ and POLε) and, in addition, LIG1, UNG2, XRCC1, DNA-dependent protein kinase (DNA PK) and cyclin A. We could not detect other factors such as cyclin B, cyclin E, PCNA, POLα or POLε (data not shown). Analysis of the composition of the Mono S fraction 15 gave perfectly comparable results, indicating that these two peak fractions represent a homogeneous preparation (data not shown).

The DNA replication, DNA repair and cell cycle proteins of the isolated complex are physically associated

In order to test whether these proteins were physically associated, we used three different approaches: co-immunoprecipitation (co-IP), native gel electrophoresis and gel filtration. We tested all our antibodies for immunoprecipitation. The best results were obtained with anti-cyclin A monoclonal antibodies, which were then used for the analysis of the peak fraction 15. As shown in Figure 4A, anti-cyclin A antibodies were able to efficiently immunoprecipitate cyclin A from the Mono S peak fraction. In addition, several proteins were found to co-immunoprecipitate together with cyclin A: POLε, XRCC1, LIG1, APE1, UNG2 and POLβ. As controls, we performed the reciprocal co-IPs for some of these interactions and we found that LIG1, cyclin A and POLε were co-IPed together with POLβ by anti-POLβ antibodies, whereas POLβ and cyclin A were co-IPed together with POLε by anti-POLε antibodies. As an additional control, POLβ-null or wild-type cells (MEFs) were lysed and subjected to immunoprecipitation with anti-POLβ antibodies. Immunoprecipitated material was analysed with anti-POLβ and anti-LIG1 antibodies. As shown in Figure 4A, right panel, only in the POLβ wild-type cells LIG1 could be co-immunoprecipitated with POLβ, whereas no immunoprecipitated proteins were detected in POLβ-null cells. The Mono S fraction was analysed by native gel electrophoresis. As shown in Figure 4B, antibodies directed against POLβ, POLε, POLζ, LIG1 and XRCC1, all reacted with a single band in the high molecular weight range (apparent Mr > 669 kDa). In order to further confirm the physical association of these proteins, the Mono S fraction 15 was subjected to gel filtration. Figure 4C shows the dot blot immunoassay of...
the eluted fractions, where it can be seen that POLα, LIG1 and cyclin A coeluted together in fractions 10–12, corresponding to an apparent molecular mass >440kDa, as expected from a multiprotein complex. We tested, in addition, the presence of another DNA replication associated protein, MCM7, and we found that it was coeluting together with POLα, LIG1 and cyclin A, indicating its association to the complex. The gel filtration fraction 10 was further analysed by western blot. As shown in Figure 4D, POLα, POLβ, POLδ, LIG1, DNA PK, cyclinA and MCM7, were detected in the same fraction.

All together, these results suggested that we have isolated a multiprotein complex containing DNA replication, DNA repair and cell cycle regulatory proteins.

Figure 2. Protein kinase activity of the multiprotein complex. (A) Phosphorylated polypeptides were revealed upon incubation of the Mono S fraction in the presence of [γ-32P] ATP alone (lane 1), or in combination with histone H1 (lane 2), histone plus p21 (lane 3) or histone plus Olomucine (lane 4). (B) The Mono S fraction was immunoprecipitated with anti-cyclin A antibodies and the immunoprecipitated material was probed with antibodies against cyclin A (lane 1) or anti-phosphoSer/Thr (lane 2).

The multiprotein complex is active in base excision repair

In order to verify the repair capacity of the replication complex an in vitro BER assay was performed by using, as substrate, a plasmid containing either an AP site or a single uracil residue. Following incubation of the plasmid with the multiprotein complex (Mono S fraction 14), the total DNA repair events (short- and long-patch repair events) were measured by [α-32P] dTTP incorporation in restriction fragment A, whereas incorporation of [α-32P] dCTP in fragment B corresponded exclusively to the long-patch repair events (see scheme, Figure 5 bottom). As shown in Figure 5 the AP site was repaired predominantly via short-patch pathway, with a fraction (25% of total) of long-patch repair events also occurring, which were largely aphidicolin-sensitive (lanes 2 and 4). Addition of PCNA increased incorporation in both A and B fragments (lanes 5 and 6) and this increase was abolished by aphidicolin (lanes 7 and 8), suggesting a role for the PCNA-dependent and aphidicolin sensitive enzymes POLδ/POLε in the repair process. Interestingly, inhibition of POLβ by specific antibodies, resulted in a switch from short- to long patch, as indicated by the increase of incorporation in fragment B (lanes 9 and 10).

When similar experiments were performed with a uracil-containing oligonucleotide (Figure 6A), the isolated complex showed uracil excision activity (lane1). This cleavage activity was due to UNG2 as demonstrated by its
Figure 4. Physical association of DNA replication and repair proteins. (A) Left panel: The Mono S fraction 15 was immunoprecipitated with antibodies against cyclin A, POL\(\beta\), POL\(\alpha\) or anonymous IgGs. Right panel: POL\(\beta\) null (\(-/-\)) or wild type (+/+)) mouse embryonic fibroblasts were lysed and the extracts used for immunoprecipitation in the presence of anti-POL\(\beta\) antibodies. The immunoprecipitated material was then immunoblotted with anti-POL\(\beta\) and anti-LIG1 antibodies, as indicated. S, supernatant (1:10); IP, immunoprecipitated material. (B) The Mono S fraction 15 was subjected to native gel electrophoresis, followed by immunoblotting analysis with antibodies against POL\(\alpha\), POL\(\beta\), POL\(\epsilon\), LIG1 and XRCC1. As marked by the asterisks, all the antibodies recognized the same high molecular weight band. (C) The Mono S fraction 15 was subjected to gel filtration. Eluted proteins were analysed by dot blot with antibodies against POL\(\alpha\), LIG1, Cyclin A and MCM7. Arrows indicate the corresponding elution points of the molecular weight markers. (D) The gel filtration fraction 10 was analysed by western blot with antibodies against POL\(\alpha\), POL\(\beta\), POL\(\delta\), DNA PK, cyclin A and MCM7.
specific inhibition by Ugi (lane 2) (13). The multiprotein complex performed BER of the uracil paired with adenine (Figure 6B) prevalently by short-patch pathway with 35% of the total repair events involving the replacement of more than one nucleotide (long-patch BER). The complex was unable to incise duplex oligonucleotides containing 8-oxo-7,8-dihydroguanine/A (8-OH-G/A) and 5-hydroxyuracil (5-OHU/A) mismatches indicating the absence of MYH and NEIL1 activities (data not shown).

The multiprotein complex is endowed with a protein kinase activity and contains two major phosphorylated species

Since the multiprotein complex isolated here contained cyclin A, which is the regulatory subunit of cyclin-dependent protein kinases (cdks), we tested whether this complex displayed cdk activity. As a reference substrate we used histone H1. As shown in Figure 2A, lane 1, incubation of the purified complex (Mono S fraction 14) in the presence of [γ-32P] ATP resulted in the appearance of two major phosphorylated polypeptides, one with an apparent MW of 110–120 kDa (high molecular weight phosphopeptide, HMP) and one with an apparent MW of 50 kDa (low molecular weight phosphopeptide, LMP). Addition of histone H1 to the reaction resulted in the appearance of an additional product at the expected position for phosphorylated H1 (lane 2). When the reaction was complemented with either the kinase inhibitor Olomucine (lane 4) or the specific cdk inhibitor protein p21 (lane 3), all phosphorylated products were greatly reduced, suggesting that they were phosphorylated by a cyclin A/cdk complex. In order to prove the physical association of the HMP and LMP proteins with the complex, a co-immunoprecipitation experiment was performed with anti-cyclin A antibodies. As shown in Figure 2B, lane 1, cyclin A was successfully immunoprecipitated. When the same sample was probed with a cocktail of antiphosphoserine–threonine antibodies, two bands appeared, with apparent MW identical to the LMP and HMP proteins detected by the kinase assay (Figure 2B, lane 2). According to its apparent electrophoretic mobility, the LMP protein appears to be distinct from cyclin A (Figure 2B, compare lane 1 with lane 2). These results seem to indicate that two major phosphopeptides are physically associated to the isolated complex and are likely phosphorylated by cyclin A/cdk. The precise identity of these two proteins is under investigation. In summary, the multiprotein complex presented here contains an active cyclinA/cdk complex which is able to phosphorylate some components of the complex itself.

DISCUSSION

The mammalian genome is constantly subjected to chemical alterations (~10^6 modifications per day) that have the potential to cause genome instability. Particular vulnerability exists during replication when attempted
replication across a damaged template can lead to dangerous lesions such as double strand breaks. To counteract these threats, cells are provided with sophisticated systems that sense DNA damage and coordinate its repair. Of particular importance for the removal of damaged bases from the DNA is the BER system (2). Although BER can be reconstituted in vitro with a few essential components, numerous studies have revealed that physical and/or functional protein–protein interactions occur at virtually every step of the BER process. These interactions involve not only the classical BER proteins, but also proteins associated with other DNA transaction pathways. These interactions could play different roles including: (i) stabilizing the associated repair protein; (ii) recruiting specific partners to the damaged site for lesion repair; (iii) altering enzymatic function or activity; and (iv) coordinating BER with other pathways of DNA metabolism. Thus, physical contact between proteins does not always result in a change in the enzymatic activity of the proteins involved, and for this reason it might even go undetected by in vitro enzymatic studies.

Here, we demonstrated the existence in human cells of a preassembled multiprotein complex containing cyclin A, DNA replication proteins and BER/SSBR components. Also one component of DSB repair, DNA PK, is present in this complex. Interestingly, the DNA PK/Ku70/Ku80 heterotrimeric complex that contributes to the resolution of DSB has been recently shown to interact with XRCC1 and stimulates its phosphorylation (14). Since we isolated this complex from nuclei of proliferating human cells without prior treatment with DNA-damaging agents, it seems that the presence of this complex is independent of damaged DNA. However, we cannot exclude that the basal physiological level of DNA damage might be enough to trigger complex formation. The isolation from undamaged cells of multiprotein BER/SSBR complexes by different methods has been previously reported (15–18). In 1996, Prasad et al. (15) reported the isolation by a glycosylase method of a complex of a molecular mass of ~180 kDa from bovine testis that contained POLβ/C24 and POLδ. In 1998, Prasad et al. (15) reported the isolation by a glycosylase method of a complex of a molecular mass of ~180 kDa from bovine testis that contained POLβ/C24 and POLδ. In 2001, Prasad et al. (15) reported the isolation by a glycosylase method of a complex of a molecular mass of ~180 kDa from bovine testis that contained POLβ/C24 and POLδ. However, in addition to UNG2, APE1, LIG1, XRCC1, POLβ and POLδ, we identified two additional DNA polymerases, namely POLε and POLδ. POLε and POLδ have been already suggested to take part in the long-patch BER (reviewed in (2)). Accordingly, our isolated complex was able to perform both short- and long-patch BER at AP sites with the short patch BER as the predominant pathway. Addition of PCNA and/or inhibition of POLβ switched the equilibrium in favour of long-patch repair products, whereas addition of aphidicolin resulted in the opposite effect. This reflects an intrinsic ability of the complex to respond to exogeneous stimuli (such as the presence of PCNA and/or limiting POLβ activity). We did not find PCNA associated to our complex. It must be stressed that we used an extraction protocol which preserves the chromatin integrity. We hypothesize that PCNA is not stably part of the complex, but rather could act as a recruiting factor for chromatin binding through physical interaction with one or more of the components of the complex.

Several BER proteins have been shown to undergo post-translational modifications. For example, FEN1, XRCC1, APE1, LIG1 and XRCC1 are phosphorylated (reviewed in (8)). Interestingly, in our complex we detected a protein kinase activity, which was inhibited by p21WAF/CIP and was able to phosphorylate endogenous polypeptides, suggesting that phosphorylation can occur within the complex itself.

The fact that the major S-phase specific cyclin A, was physically associated to the complex together with the major replicative enzyme POLε and with the DNA replication protein MCM7 immediately suggests a possible link of the BER/SSBR pathway with the DNA replication process. Recent data indicated that the DSB pathway is controlled by the cyclin A–cdk complex (23) suggesting a possible model of co-regulation of repair pathways during the cell cycle (24). A preassembled complex containing DNA replication, cell cycle and BER components specifically involved in endogenous DNA damage processing (i.e. UNG2 and APE1) might indicate the need of a repairosome for these lesions frequently encountered by the replication machinery. Indeed, UNG2 has been found associated to DNA replication proteins (25–27). The present data, however, does not allow to conclude that the isolated repairosome is specifically formed in S-phase or exists also in other phases of the cell cycle, such as G1 or G2. Analysis of the dynamics of the repairosome as a function of the cell cycle is currently underway in our laboratory. Interestingly, the DSB repair enzyme DNAPK is also present in our BER complex, along with XRCC1. XRCC1 has been shown to coordinate the assembly of SSBR and BER components at damaged sites and is essential for cells survival after SSB induction (reviewed in (28)). In addition, recent results suggest that XRCC1 phosphorylation by DNA PK in response to ionizing radiation, might trigger a signal cascade leading to NHEJ-dependent DSBR (14).
Thus, XRCC1 appears to act as an early responder of DNA breaks at stalled replication forks. In light of these results, our findings raise the intriguing hypothesis that XRCC1 and DNA-PK might be stably associated to a repairosome, which is linked to the DNA replication fork through association with POLα and MCM7. Depending on the cellular context, cell cycle phase and type of damage, this repairosome will have the ability to quickly respond to a stalled replication fork, activating different DNA repair pathways.

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REFERENCES