Cytosolic PCNA interacts with p47phox and controls NADPH oxidase NOX2 activation in neutrophils

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Neutrophils produce high levels of reactive oxygen species (ROS) by NADPH oxidase that are crucial for host defense but can lead to tissue injury when produced in excess. We previously described that proliferating cell nuclear antigen (PCNA), a nuclear scaffolding protein pivotal in DNA synthesis, controls neutrophil survival through its cytosolic association with procaspases. We herein showed that PCNA associated with p47phox, a key subunit of NADPH oxidase, and that this association regulated ROS production. Surface plasmon resonance and crystallography techniques demonstrated that the interdomain-connecting loop of PCNA interacted directly with the phox homology (PX) domain of the p47phox. PCNA inhibition by competing peptides or by T2AA, a small-molecule PCNA inhibitor, decreased NADPH oxidase activation in vitro. Furthermore, T2AA provided a therapeutic benefit in mice during trinitro-benzene-sulfonic acid (TNBS)-induced colitis by decreasing oxidative stress, accelerating mucosal repair, and promoting the resolution of inflammation. Our data suggest that targeting PCNA in inflammatory neutrophils holds promise as a multifaceted antiinflammatory strategy.

Introduction

Neutrophils are key cells of the innate immune system (Mantovani et al., 2011) but may also be involved in pathological inflammatory processes when their activation and elimination are not properly regulated (Soehnlein et al., 2017). At sites of inflammation, neutrophils release biological molecules including proteases and ROS that can kill pathogens but also cause host tissue damage (Nauseef, 2007). The enzyme responsible for ROS production in neutrophils is the phagocyte NADPH oxidase (NOX2), a complex composed of two membrane proteins (gp91phox, p22phox) and four cytosolic proteins (p47phox, p67phox, p40phox, and the small GTPase Rac1/2) that assemble upon activation (El-Benna et al., 2016). Although ROS production is important for innate immunity, excessive ROS release can induce oxidative stress, leading to cell death and tissue injury (Nathan and Cunningham-Bussel, 2013). ROS are involved in numerous inflammatory diseases such as rheumatoid arthritis, inflammatory bowel disease, atherosclerosis, and aging (Lambeth et al., 2008). Coordination between neutrophil activation, ROS production, and cell death is pivotal in both host defense and the control of inflammation (Kennedy and DeLeo, 2009; Geering and Simon, 2011).

In this context, we have previously shown that the proliferating cell nuclear antigen (PCNA) protein plays a key role in neutrophil apoptosis (Witko-Sarsat et al., 2010; Ohayon et al., 2016). In the majority of cell types, PCNA is an exclusively nuclear protein that regulates processes related to DNA replication including cell cycle regulation and recombination (Maga and Hubscher, 2003). It has been dubbed the “maestro of the replication fork” (Moldovan et al., 2007). PCNA exists as a homotrimer that encircles duplex DNA, forming a ring-shaped clamp. PCNA is a trimer, and each monomer is composed of two similarly folded globular regions linked by a flexible interdomain-connecting loop, which is the preferred site for the binding of partner molecules (Warbrick, 1998). PCNA is a scaffolding
protein and has no intrinsic enzymatic activity. Reflecting the wide variety of roles of PCNA, >100 PCNA-interacting proteins involved in DNA-related processes have been identified (Maga and Hubscher, 2003).

In mature neutrophils, on the other hand, PCNA is expressed exclusively in the cytosol. PCNA is exported from the nucleus at the end of granulocytic differentiation via a nuclear export sequence we have previously identified (Bouayad et al., 2012). PCNA dictates neutrophil survival by binding to apoptotic procaspases, sequestering and preventing their activation (Witko-Sarsat et al., 2010). Neutrophils are differently differentiated nonproliferative cells, suggesting the cytosolic PCNA may coordinate functions apart from those related to proliferation. Cytosolic PCNA also promotes survival in acute myeloid leukemic cells resistant to chemotherapy. Furthermore, PCNA promotes glycolysis in these cells, serving as a scaffold for glycolytic proteins (Ohayon et al., 2016), suggesting the cytosolic PCNA may coordinate functions apart from those related to proliferation. Cytosolic PCNA also promotes survival in acute myeloid leukemic cells resistant to chemotherapy. Whether PCNA has other binding partners in the cytosol of neutrophils and what contribution PCNA makes to neutrophil function more broadly remains poorly understood (Witko-Sarsat and Ohayon, 2016).

Given these roles of PCNA as a scaffold, we performed an exploratory proteomic analysis testing for novel PCNA partners within the neutrophil cytosol. We show using a number of techniques that PCNA associates with p47phox, a key subunit of the NADPH oxidase complex, and that this association regulates ROS production. Interfering with this association resulted in a potent in vivo antiinflammatory effect in two neutrophil-dominated models of inflammation, zymosan-induced peritonitis and trinitro-benzene-sulfonic acid (TNBS)–induced colitis.

Results

Cytosolic PCNA interacts with the phox homology (PX) domain of p47phox

We identified PCNA partners by communoprecipitation and mass spectrometry (MS) analysis. PCNA was associated with proteins involved in cytoskeleton function, oxidant regulation, cell signaling, and glycolysis, as previously described in HL-60 cells (Ohayon et al., 2016; Table 1). Notably, the cytosolic components of the NADPH oxidase/NOX2, namely p47phox, p67phox, and p40phox as well as kinases and GTP-binding proteins, were associated with PCNA, suggesting a potential role of PCNA in the modulation of NADPH oxidase activity. As p47phox, p67phox, and p40phox form a complex in the cytosol, we examined the individual in vitro interactions between recombinant PCNA and p47phox, p67phox, p40phox, and Rac2 by surface plasmon resonance (SPR). A strong interaction was observed between p47phox and PCNA in a dose-dependent manner, with an equilibrium dissociation constant ($K_D$) value within the nanomolar range ($K_D = 7.7 \times 10^{-8}$ M; Fig. 1 A and Table 2). No direct interaction was detected between PCNA and p67phox or p40phox, whereas Rac2 could interact with PCNA with a micromolar-range affinity. To identify the domain of p47phox involved in PCNA interaction, we first examined if this interaction could be prevented by phosphoinositides that can bind to the PX domain (residues 1–141) that is crucial for p47phox membrane association (Kanai et al., 2001). When phosphoinositides and p47phox were preincubated before injection over immobilized PCNA, the percentage of p47phox binding to PCNA was decreased in a dose-dependent manner (Fig. 1 B). In keeping with this hypothesis, recombinant PX domain generated as a truncated p47phox protein showed a direct interaction with PCNA, although the calculated $K_D$ was higher than that of full-length p47phox. This suggests that either p47phox binding is dependent on the conformation of the PX domain in intact p47phox or other p47phox domains are involved in PCNA binding (Fig. 1 C and Table 2).

Many PCNA-interacting proteins including p21/waf1 contain a conserved sequence known as PCNA-interacting protein box (PIP-box) with a consensus sequence of [Q|x][h][XX][a][a], where h stands for hydrophobic and a for aromatic residues (Fig. S1). We identified four peptide sequences (p47phox-46–67, p47phox-63–90, p47phox-83–110, and p47phox-106–127) containing a PIP-box motif that were not buried in p47phox PX domain (Fig. 1 D) sequence. The corresponding p47phox peptides were tested for their binding capacity to PCNA by SPR and were compared with the p21-141-160-peptide used as a positive control ($K_D = 10^{-7}$ M) and to the mutant p21 peptide deprived of PCNA binding sites as a negative control (Zheleva et al., 2000; Warbrick, 2006). Peptides p47phox-46–67, p47phox-63–90, and p47phox-83–110 did not interact with PCNA, while p47phox-106–127 showed a direct interaction with PCNA (Fig. 1 E and Table 2). Interestingly, the p47phox-106–127 peptide and the p47phox PX domain bind PCNA with a similar affinity ($K_D = 5 \times 10^{-5}$ M; Table 2), strongly suggesting that the p47phox-106–127 sequence was mainly responsible for the p47phox PX–PCNA interaction.

Coinjection of p21 peptide (2, 5, or 8 µM) with the p47phox over immobilized PCNA decreased the binding of p47phox to PCNA in a dose-dependent manner, compared with binding in the absence of the p21 peptide (Fig. 1 F). Likewise, similarly to the p21 peptide and in agreement with its ability to bind PCNA, the p47phox-106–127 peptide inhibited the interaction between PCNA and the p47phox protein in a dose-dependent manner, as measured by SPR. These data unambiguously demonstrated that the interdomain-connecting loop of PCNA was involved in the interaction with p47phox and strongly suggested that this interaction was mediated by the p47phox PX domain mapping the 106–127 residues.

To further ascertain the molecular association between PCNA and p47phox-106–127 peptide, crystallization assays and diffraction analysis were used. Crystals could be grown under several different conditions, but one (see Materials and methods for details) provided reproducible crystals diffracting at ~3-Å resolution. The unit cell dimensions and the crystal packing were similar for all crystals (Figs. 2 A and SI, B–D). However, the space group was found to be either orthorhombic (P2_12_1_2, one trimer in the asymmetric unit) or monoclinic (P2_1, two trimers in the asymmetric unit). Our attempts to cocrystallize intact p47phox or its PX domain together with PCNA were unsuccessful and were hampered by the flexible p47phox structure. However, we could obtain crystallographic data by soaking PCNA crystals in solutions containing 100–500 µM of p47phox-106–127 peptide for 24 h before data collection. Complete
crystallographic data were collected on ∼50 crystals, with a maximum resolution varying from 4.2 to 2.8 Å, and the PCNA structure was determined for all these crystals by molecular replacement using the PCNA 1AXC structure. In five crystals, we observed a significant positive electron density on the residual \((F_{\text{obs}} - F_{\text{calc}})\) electron density map, indicating that the p47phox-106-127 peptide could bind the PCNA interdomain-connecting loop region (Fig. 2 B).

Although the maximum resolution (3.22 Å) and the occupancy of the bound peptide obtained for these five crystals was

| Swiss-Prot ID | Protein name | Molecular weight | Gene name | UniProt ID |
|---------------|--------------|------------------|-----------|------------|
| UBA1_HUMAN    | Ubiquitin-like modifier-activating enzyme 1 | 117.8 | UBA1 | P22314 |
| PROF1_HUMAN   | Profilin-1   | 15.1             | PFN1      | P07737 |
| FLNA_HUMAN    | Filamin-A    | 280.7            | FLNA      | P21333 |
| MOES_HUMAN    | Moesin       | 67.8             | MSN       | P26038 |
| VINC_HUMAN    | Vinculin     | 123.8            | VCL       | P18206 |
| ARCI1B_HUMAN  | Actin-related protein 2/3 complex subunit 1B | 40.9 | ARPC1B | O15143 |
| PDIA1_HUMAN   | Protein disulfide-isomerase              | 57.1 | P4HB | P07237 |
| THIO_HUMAN    | Thioredoxin                                      | 11.7 | TXN | P10599 |
| IQGAP1_HUMAN  | Ras GTPase-activating-like protein IQGAP1       | 189.3 | IQGAP1 | P46940 |
| GD2B_HUMAN    | Rab GDP dissociation inhibitor β              | 50.7  | GD2 | P50195 |
| GDI1_HUMAN    | Rho GDP-dissociation inhibitor 1              | 23.2  | ARHGDI1 | P52565 |
| GDI2_HUMAN    | Rho GDP-dissociation inhibitor 2              | 23    | ARHGDI2 | P52566 |
| NCF1_HUMAN    | Neutrophil cytosol factor 1; p47phox         | 44.8  | NCF1 | P14598 |
| NCF2_HUMAN    | Neutrophil cytosol factor 2; p67phox         | 59.8  | NCF2 | P19878 |
| NCF4_HUMAN    | Neutrophil cytosol factor 4; p40phox         | 39    | NCF4 | Q15080 |
| COR1A_HUMAN   | Coronin-1A                                        | 51    | CORO1A | P31146 |
| GNAI2_HUMAN   | Guanine nucleotide-binding protein G(i) subunit α-2 | 40.5 | GNAI2 | P04899 |
| G3P_HUMAN     | Glyceraldehyde-3-phosphate dehydrogenase       | 36.1  | GAPDH | P04406 |
| G6PD_HUMAN    | Glucose-6-phosphate 1-dehydrogenase           | 59.3  | G6PD | P11413 |
| G6PL_HUMAN    | Glucose-6-phosphate isomerase                 | 63.1  | GPI  | P06744 |
| 6PGD_HUMAN    | 6-Phosphogluconate dehydrogenase              | 53.1  | PGD  | P52209 |
| 6PGL_HUMAN    | 6-Phosphogluconolactonase                     | 27.5  | PGLS | Q95336 |
| ALDOA_HUMAN   | Fructose-bisphosphate aldolase A              | 39.4  | ALDOA | P04075 |
| ALDOC_HUMAN   | Fructose-bisphosphate aldolase C              | 39.5  | ALDOC | P09972 |
| HK1_HUMAN     | Hexokinase-1                                   | 102.5 | HK1  | P19367 |
| LDHAL6B_HUMAN | L-lactate dehydrogenase A-like 6B             | 41.9  | LDHAL6B | Q98Y2Z |
| LDHA_HUMAN    | L-lactate dehydrogenase A chain               | 36.7  | LDHA | P00338 |
| LDHB_HUMAN    | L-lactate dehydrogenase B chain               | 36.6  | LDHB | P07195 |
| PKM_HUMAN     | Pyruvate kinase isozymes M1/M2                | 57.9  | PKM  | P14618 |

Immunoprecipitation of PCNA was performed in cytosols from human neutrophils obtained by nitrogen cavitation, and proteins associated with PCNA were identified by MS. This representative experiment was performed two times on two different cytosols with similar results.
too low to determine unambiguously the amino acids of p47phox-106-127 involved in the interaction with PCNA, we propose two computational models of the complex between PCNA and p47phox PX domain, based on structures available in the Protein Data Bank (PDB). We used the structure of the complex between the p21 peptide (mapping the 141–160 residues) and PCNA (PDB entry 1AXC; Gulbis et al., 1996) and the structure of the p47phox PX domain (PDB entry 1KQ6). In the first option, structural analysis of the 1KQ6 structure showed that the 114–118 PIP-box of p47phox was partially buried and should adopt another conformation to be able to interact with PCNA. Therefore, the p47phox PX-106-127 region has been modeled to adopt a conformation close to the one adopted by p21 in 1AXC structure (Fig. 2 C). If such a mode of interaction is plausible, as it generates no steric hindrance, it relies on a hypothetical conformational change in the 106–127 region of the p47phox PX domain that remains to be experimentally observed. The second option was to search for regions of the p47phox that adopt a conformation that is similar to the one adopted by the p21 peptide, when bound to PCNA. The 122–126 region of the p47phox PX domain is structurally very similar to the 147–151 region of p21 (root mean square difference of 0.25 Å) and corresponds to the core of the PIP-box that was structurally conserved in numerous PCNA-peptide complex structures. By superimposing the 122–126 residues of 1KQ6 p47phox structure on the 147–151 region of 1AXC PCNA/p21 peptide, we built a plausible PCNA-p47phox PX domain complex with no steric clashes (Fig. 2 D). The distinction between these two possibilities should await the crystal structure of p47phox protein, which is not yet publicly available.

PCNA is involved in NOX2 activation in intact cells
To address the functional impact of the PCNA–NOX2 interaction, we first used neutrophil-like differentiated PLB985 cells, an established model of human neutrophils, which allow transfection experiments. We previously reported that in proliferating nondifferentiated PLB985 cells, PCNA is nuclear (Fig. 3 A) but becomes exclusively cytosolic after differentiation, therefore representing a suitable surrogate for human neutrophils (De Chiara et al., 2013). Differentiated PLB985 cells that stably overexpress PCNA as previously described (Witko-Sarsat et al., 2010) showed a marked increase in PMN- and opsonized zymosan (OZ)-induced superoxide production compared with PLB985 cells stably transfected with control plasmid (Fig. 3, B...
and C). Increased PMA-induced ROS production in PCNA-overexpressing cells was confirmed by 2',7'-dichlorofluorescin diacetate (DCF-DA) FACS analysis. Notably, this increase in ROS production was abolished when PLB985 cells were transfected with PCNA fused to the SV40 nuclear locali-
sation sequence (NLS), thereby excluding from the cytosol (Bouayad et al., 2012; Fig. 3, D and E). Transfection of PCNA siRNA efficiently down-regulated PCNA protein expression (Fig. 3 F) and significantly decreased ROS production measured by luminol chemiluminescence (CL) in response to PMA and OZ (Fig. 3, G and H). Likewise, we used dimethylformamide (DMF)-differentiated PLB985 cells overexpressing the protein p21/waf1 (Fig. 3 I), which can bind PCNA and inhibit its antiapoptotic function (Martin et al., 2016). DMF-differentiated PLB985-p21A45R cells have significantly decreased NADPH oxidase activity triggered by either PMA or OZ relative to the PLB985 cells expressing an empty plasmid (Fig. 3, J and K). Notably, the effect of inhibiting PCNA on NADPH oxidase function is not merely a consequence of loss of neutrophil viability. The decreased NADPH oxidase function in differentiated PLB985 cells by PCNA inhibition (by either siRNA or ectopic expression of p21/waf1) was evident at time points in the absence of significant cell death (Fig. S2, A and B). Altogether, these results demonstrated that cytosolic PCNA controlled NADPH oxidase-dependent ROS production in intact cells.

Inhibition of PCNA–p47phox interaction hindered NADPH oxidase activation

We next examined if targeting PCNA could affect NADPH oxidase assembly in a cell-free system consisting of the cytosolic fraction containing PCNA and the NADPH oxidase components (p47phox, p67phox, p40phox, rac2) and the membrane fraction containing gp91phox and p22phox isolated from resting human neutrophils. NADPH oxidase activation was triggered by Li-SDS and the production of superoxide anions was measured by the cytochrome c reduction assay. The specificity of this assay was validated by the inhibitory effect of superoxide dismutase (SOD) or diphenylene iodonium (DPI), a NADPH oxidase inhibitor (Fig. 4 A). In this cell-free system, we tested the effect of the p21 peptide (Fig. 4 B) and T2 amino alcohol (T2AA), a nonpeptide cell-permeable PCNA inhibitor (Punchihewa et al., 2012; Inoue et al., 2014). Both compounds bind specifically to the PCNA interdomain-connecting loop to inhibit interactions with its partners (Fig. 4 C). Each compound led to a dose-dependent decrease in superoxide anion production, thereby demonstrating that PCNA was involved in NOX2 activation. T2AA also decreased PMA-, formyl-methionyl-leucyl phenylalanine (f-MLF)–, and OZ-induced NADPH oxidase activation in intact neutrophils measured by luminol CL in a dose-dependent manner (Fig. 4, D–F). The inhibitory effect of T2AA on ROS production was not related to apoptosis induction, since no Annexin-V labeling was detected on neutrophils after 1-h T2AA treatment up to a concentration of 50 µM in the presence of PMA (Fig. S2, C and D). Moreover, inhibition of caspases by the pan-caspase inhibitor Z-VAD(OMe)-FMK (Z-VAD) did not abrogate the inhibitory effect of T2AA on f-MLF-induced respiratory burst (Fig. S2, E and F). To further characterize the interaction between PCNA and p47phox, we examined whether PCNA, like cytosolic NADPH oxidase components, could translocate to the membrane upon PMA activation. Western blot analysis of neutrophil membranes and cytosol clearly showed that both p47phox and PCNA translocated to the membrane fraction upon PMA activation.
Notably, T2AA inhibited p47phox translocation to the membrane during activation without inhibiting its phosphorylation. Consistent with a direct interaction between p47phox and PCNA, we observed that PCNA and p47phox colocalized in cytosol in unstimulated neutrophils by confocal microscopy analysis after indirect immunofluorescence labeling (Fig. 5 A). Using Duolink technology, a proximity ligation assay, we provided evidence that PCNA and p47phox are in close proximity within the cytosol of resting neutrophils (Fig. 5 B). In contrast, p47phox and rac2, two cytosolic proteins known to localize in separate cytosolic complexes in resting neutrophils, did not demonstrate proximity, supporting the specificity of this technology. Interestingly, PCNA and p47phox proximity was strongly decreased after PMA activation. Notably, immunoprecipitation of PCNA in the cytosol of resting neutrophils showed that it coimmunoprecipitated with p47phox (Fig. 5 C). However, the amount of p47phox associated with PCNA was strongly decreased after PMA or f-MLF stimulation. Taken together, our results suggested that PCNA associates with p47phox in cytosol of resting neutrophils and facilitates NADPH oxidase assembly. The association is disrupted after activation, although PCNA could translocate to the plasma membrane. Importantly, T2AA did not inhibit other antipathogen functions such as f-MLF-induced degranulation, C5a- or f-MLF-induced chemotaxis, and *Escherichia coli* phagocytosis tested in vitro on isolated neutrophils (Fig. S3).

Destabilization of PCNA scaffold by T2AA inhibits NADPH oxidase activity in vivo

We next evaluated whether inhibiting the interaction between PCNA and p47phox using T2AA would inhibit NADPH oxidase activity in vivo in inflammatory neutrophils during zymosan-induced peritonitis in mice (Martin et al., 2016). Peritoneal lavages were collected 4 h after zymosan injection, and inflammatory cells including neutrophils and macrophages were analyzed by FACS. T2AA did not have any significant effect on the total number of cells (Fig. 6 A) or the cellular composition in the lavage fluid (Fig. 6, B and C). Peritoneal neutrophils from T2AA-treated mice showed decreased ex vivo ROS production compared with controls (Fig. 6 D), thereby confirming our in vitro results on human neutrophils (Fig. 4 D). We next used in vivo noninvasive imaging of neutrophil activation using L012-enhanced CL (Kielland et al., 2009) to demonstrate that T2AA could inhibit NADPH oxidase in situ. Zymosan-induced peritonitis increased L012-CL in the peritoneum, and T2AA significantly attenuated this (Fig. 6, E and F). In our previous study, we had shown that p21/waf1 protein acts as a specific inhibitor of the PCNA scaffold in inflammatory neutrophils (Martin et al., 2016). We further investigated NADPH oxidase activation in inflammatory neutrophils in p21/waf1−/− mice. PMA-induced ROS production was significantly increased in peritoneal lavage neutrophils from p21/waf1−/− mice compared with neutrophils from WT mice, confirming that binding of p21/waf1 to PCNA restraints NADPH oxidase activation (Fig. 6, G and H).

**Figure 2.** Electron density map of PCNA crystals associated with the p47phox-106-127 peptide and molecular modeling of the complex PCNA-p47phox PX domain. (A) Diffraction pattern of PCNA crystals soaked in p47phox-106-127 peptide solution and snapshot of the frozen crystal. This experiment was performed three times with identical results. (B) Residual \( (F_{\text{obs}} - F_{\text{calc}}) \) electron density map from 3.23-Å-resolution crystallographic data. The three highest peaks of the residual \( (F_{\text{obs}} - F_{\text{calc}}) \) electron density map (green) are located where PIP-box-containing peptides interact with PCNA. The p21-139-160 peptide from PCNA-p21-139-160 peptide complex structure (PDB entry 1AXC) in cyan with the four main residues of the PIP-box highlighted in red. (C and D) Two putative models of PCNA-p47phox-PX domain complex based on the crystal structures of human PCNA (blue) the structure of p21-139-160 peptide (cyan) bound to PCNA and the p47phox PX domain (purple) with the residues 106–127 (gold). Predicted PIP-box residues are depicted in red. (C) Model based on the putative PIP-box of the p47phox-106-127 peptide whose conformation has been modified to fit the p21 peptide structure. (D) Model based on the structural homology between the helix turn of p47phox-122-126 and of the p21-PIP-box.
Figure 3. PCNA controls ROS production in differentiated PLB985 cells. (A) Representative experiment out of 10 showing the detection of cytosolic PCNA by immunofluorescence in DMF-differentiated PLB985 cells. The scale bars correspond to 10 µm, magnification 63×. (B) Kinetic analysis of luminol-CL in DMF-differentiated PLB985 cells transfected with a control (pCT) or with a PCNA-coding plasmid stimulated or not (resting) with PMA or OZ. (C) CL peaks corresponding to B were expressed as mean ± SEM of four independent experiments performed in duplicate. (D) FACS plots of DCF-mediated fluorescence in PLB985-overexpressing PCNA or a nuclear form of PCNA (PCNA-NLS) versus control (pCT). Representative experiment before (-) and after (+) PMA out of four sets of independent experiments. (E) Quantification of DCF-MFI. Data are mean ± SEM (n = 10 for PCNA WT and CT, n = 4 for PCNA NLS). (F–H) Effect of PCNA siRNA on NADPH oxidase activation. (F) Western blot analysis of PCNA and actin expression after siRNA-PCNA treatment compared with siRNA-CT (upper panel) and quantification of PCNA/β-actin ratio expressed as mean ± SEM of four independent experiments performed in duplicate (lower panel). **, P < 0.01, Mann–Whitney U test. (G) Kinetic analysis of luminol-CL of DMF-differentiated PLB985 cells stimulated with PMA or OZ compared with resting. (H) CL peaks corresponding to G expressed as mean ± SEM of four independent experiments performed in duplicate. (I–K) Effect of p21/Waf1 expression on NADPH oxidase activation. (I) Western blot analysis of PCNA, p21/Waf1, PCNA, and actin expression (upper panel) and quantification of PCNA/p21/Waf1 ratio expressed as mean ± SEM of three independent experiments performed in duplicate (lower panel). *, P < 0.01, Mann–Whitney U test. (J) Kinetic analysis of luminol-CL of DMF-differentiated PLB985 cells transfected with a p21/Waf1-coding plasmid or a control plasmid (pCT) and stimulated with PMA or OZ. (K) CL peaks corresponding to J expressed as mean ± SEM of eight independent experiments performed in duplicate. ANOVA test was performed (C, E, H, and K). *, P < 0.05; **, P < 0.01; ***, P < 0.001.
As we previously reported that PCNA inhibition by the p21 peptide promotes neutrophil apoptosis, we examined whether T2AA could also affect neutrophil survival. Indeed, T2AA did not induce ex vivo apoptosis in resting bone marrow neutrophils but induced dose-dependent apoptosis in inflammatory neutrophils isolated from peritoneal lavages (Fig. S4 A). Analysis of neutrophils from WT and p47phox−/− mice (Kasahara et al., 1997) showed that this proapoptotic effect of T2AA on inflammatory neutrophils was not dependent on the presence of p47phox (Fig. S4 B). From this set of data, we conclude that T2AA triggered apoptosis preferentially in neutrophils at the inflammatory site without affecting their survival in bone marrow.

T2AA exerts a potent antiinflammatory effect in TNBS-induced colitis

As T2AA inhibited ROS production in inflammatory neutrophils, we investigated whether it could exert an antiinflammatory effect in TNBS-induced colitis (Wallace et al., 1989), as this...
model is dependent on early massive recruitment of neutrophils (Krimi et al., 2008). Mice received two doses of T2AA or vehicle by intraperitoneal injection after TNBS treatment. The increase in colonic weight/length ratio induced by TNBS, a macroscopic indicator of inflammation, was significantly decreased in the presence of T2AA (Fig. 7 A). These data were supported by the lower Wallace score in T2AA-treated mice, reflecting less extensive inflammatory lesions, with a mean percentage of protection of 50.9 ± 8.8% (Fig. 7 B). Histological analysis confirmed that TNBS induced strong acute colonic inflammation and necrosis, with extensive ulceration in untreated animals (Fig. 7 D). The colons of T2AA-treated mice displayed less ulceration and more advanced healing of the eroded surface. Reepithelialization and regeneration of colonic mucosa were illustrated respectively by colonization of the ulcer surface by epithelial cells and by mucosal repair of neighboring ulceration with gland hyperplasia and numerous mitotic cells. The T2AA beneficial effect was demonstrated by reduction in the Ameho score (Ameho et al., 1997), reflecting less-severe histological tissue damage (Fig. 7 D). We also evaluated in vivo ROS-mediated tissue damage by measuring malondialdehyde (MDA), a product of lipid peroxidation, in colon homogenates (Fig. 7 E). Our results show a significant decrease in MDA concentrations in tissues from T2AA-treated mice compared with untreated mice, in agreement with decreased MPO content. Using the in vivo L012-based imaging method, we confirmed that neutrophil NADPH oxidase was activated in TNBS-induced colitis and that T2AA significantly decreased L012-CL, indicative of a neutrophil-targeted anti-inflammatory effect (Fig. 7, F and G). Finally, neutrophil depletion was used to further ascertain that T2AA targeted activated inflammatory neutrophils. Neutrophil depletion using anti-Ly6G mAb (Coffelt et al., 2015; Fig. 8 A) resulted in a drastic loss of neutrophils in both blood and bone marrow compared with control IgG-treated mice (Fig. 8 B). Neutrophil depletion induced a significant decreased severity of colitis in the absence of T2AA as evidenced by the macroscopic analysis quantified by the Wallace score (Fig. 8 C). Notably, while T2AA exerted a
significant antiinflammatory effect in IgG-treated mice, no effect of T2AA was observed in anti-Ly6G–treated mice. Histological evaluation and immunolabeling of neutrophils confirmed the strong decrease in neutrophils in the colon of anti-Ly6G-depleted and TNBS-treated mice (Fig. 8 E). Importantly, no beneficial effect of T2AA on colon inflammation assessed by neutrophil immunolabeling was observed in neutrophil-depleted mice, albeit with a lower baseline level of disease activity (Fig. 8 E). These new data show that the acute inflammation observed in TNBS-induced colitis as well as the antiinflammatory effect of T2AA are both dependent on the presence of neutrophils.

Discussion
To effectively limit neutrophil-mediated tissue inflammation and damage, therapeutic strategies should ideally target neutrophil release of toxic mediators such as excessive ROS while promoting neutrophil apoptosis and subsequent clearance. Therapies should avoid excessive impairment of neutrophil bone marrow production to limit immunosuppressive consequences. In the current study, we have uncovered a new role of cytosolic PCNA in the regulation of NADPH oxidase activity and highlighted the therapeutic potential of targeting PCNA.

The spatial separation of the NADPH oxidase components between cytosol and membrane and their interaction in complexes ensures that the enzyme is dormant in resting cells (Boussetta et al., 2010). In this setting, PCNA binds to the p47phox within the cytosol to restrain NADPH oxidase assembly and activation. Conversely, upon stimulation, PCNA bound to p47phox translocates to the membrane to facilitate NADPH assembly. Inhibiting PCNA prevents translocation without interfering with p47phox phosphorylation, thereby acting as a novel and unexpected regulator of neutrophil activation. It is noteworthy that, while almost all p67phox and p40phox are in a complex together with p47phox, excess p47phox can be found free of the other cytosolic proteins (Iyer et al., 1994; Park et al., 1994). This unbound p47phox has been shown to interact with...
other cytosolic proteins including protein kinase C (Reeves et al., 1999) and phospholipase A2 (Shmelzer et al., 2008) and may be associated with PCNA. Other proteins identified as PCNA partners include cytoskeletal proteins, suggesting that this complex can facilitate the assembly of active NADPH oxidase complex at the membrane. Moreover, PCNA association with glycolysis enzymes, as previously shown in leukemic cells resistant to chemotherapy (Ohayon et al., 2016), strongly suggests that PCNA could be a link between survival, ROS production, and metabolism (Baillet et al., 2017).

We further demonstrated that PCNA binds to the PX domain of p47phox, in particular the region mapping the 106-127 residues enzymes, as previously shown in leukemic cells resistant to chemotherapy (Ohayon et al., 2016), strongly suggests that PCNA could be a link between survival, ROS production, and metabolism (Baillet et al., 2017).

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Figure 7. Antiinflammatory effect of T2AA on TNBS-induced colitis. (A and B) Macroscopic changes following TNBS treatment are represented by colon weight/length ratio (A) and Wallace score (B) in the respective histograms. (C) Ameho score representative of histological lesions is shown in the histogram. The data (A–C) are mean ± SEM (**, P < 0.01; ***, P < 0.001, ANOVA) from six to eight mice per group in two independent experiments. (D) Photomicrographs are representative of H&E-stained slides (upper panels) and of MPO immunohistochemical labeling (lower panels) of paraffin-embedded colonic tissues recovered from controls, TNBS alone, and T2AA. The scale bars correspond to 20 µm, magnification 200×. RG, regeneration; RE, reepithelization. (E) Colorimetric measurement of MDA in colon. The data are mean ± SEM (n = 6), **, P < 0.01, ANOVA. (F and G) Inhibitory effect of T2AA on in vivo ROS production measured by L012-CL in TNBS-induced colitis mice. (F) Representative imaging of ROS production in individual mice with untreated TNBS-induced colitis (right panel) or T2AA-treated TNBS-induced colitis (left panel). This representative experiment shows the pseudocolor heat map representing photons/s/cm²/steradian. (G) Quantification of in vivo L012-CL from three to four mice per group using the percentages of L012-CL variation between 24 and 48 h. The data are expressed as mean ± SEM (*, P < 0.05, nonparametric Wilcoxon test).
involved in lipid binding. This is consistent with the notion that PCNA inhibits p47phox binding to the membrane in the resting state. Indeed the crystal structure of the isolated PX domain simultaneously bound to phosphatidylinositol 3,4-bisphosphate and phosphatidic acid revealed that the two lipids bind to two basic pockets (Karathanassis et al., 2002; El-Benna et al., 2009). This study further showed that a point mutation of the second SH3 domain or p47phox phosphorylation induced a transition from a closed to an open conformation that binds membranes through the PX domain (Marcoux et al., 2010). Our structural analysis using both crystallography and SPR techniques allows us to propose two plausible molecular models for the physical association between the interdomain-connecting loop domain of PCNA and the PX domain of p47phox. Interestingly, the p47phox 106–127 peptide that we have identified as the PCNA-interacting site in the PX domain has been previously reported to strongly inhibit NADPH oxidase activation in a cell-free system by using a peptide-walking technique to map p47phox functional domains (Morozov et al., 1998), confirming that this p47phox sequence indeed has a functional role in the control of NOX2 activation. Because PCNA is a trimer, it can potentially bind up to three PIP-motif-containing proteins at the same time (Boehm and Washington, 2016). Although the interdomain-connecting loop of PCNA can bind numerous PIP-box-containing proteins, other domains of PCNA are involved in the binding with partner proteins including p47phox, thereby increasing the diversity of interacting partners that PCNA can accommodate to orchestrate complex biological processes.

We have previously reported that the PCNA scaffold is dynamic and is adapted to the inflammatory state of the cell (Martin et al., 2016). Indeed, PCNA binding to procaspases and to p47phox has important consequences for cell survival and NADPH oxidase activation, respectively. Pertinently, p47phox-PCNA association by maintaining NOX2 in a resting state contributes to neutrophil survival. Nonetheless, PCNA–p47phox interaction is not essential for PCNA anti-apoptotic activity, since T2AA triggers apoptosis in inflammatory neutrophils in both WT and p47phox−/− mice.

Our study demonstrated for the first time that interfering with PCNA scaffold could be the basis of antioxidant and proresolution therapy in neutrophil-induced inflammation. A hallmark of

Figure 8. Absence of T2AA antiinflammatory effect in TNBS-induced colitis following neutrophil depletion. (A) Experimental protocol of colitis after neutrophil depletion using anti-Ly6G mAb. (B and C) Flow cytometry quantification of neutrophils in blood (left panel) and in bone marrow (right panel) in control (IgG) and neutrophil-depleted (Ly6G) mice. (C) Macroscopic changes following TNBS treatment measured by Wallace score. The data are mean ± SEM (ANOVA). (D) Representative photomicrographs of neutrophil immunostaining of colonic tissues with the neutrophil marker recovered from control (IgG) and neutrophil-depleted mice (Ly6G) treated with TNBS alone or TNBS combined with T2AA. The scale bars correspond to 20 µm, magnification 200×. (E) Quantification of neutrophil immunostaining as described in Materials and methods. In B, C, and E, the data are mean ± SEM. Two independent sets of experiments were performed with a total of eight mice per group. *, P < 0.05; **, P < 0.01; ***, P < 0.001, ANOVA.
inflammatory bowel diseases including Crohn’s disease and ulcerative colitis is excessive neutrophil accumulation in the colon, resulting in the release of toxic proteases and ROS (Parkos, 2016). Excess neutrophil-derived ROS participate in tissue damage in several inflammatory diseases, so inhibition of NADPH oxidase activation could represent a relevant therapeutic strategy (Lambeth et al., 2008).

The role of ROS in inflammation is complex, as they play a role in tissue damage as well as a positive role in the resolution of inflammation (Nathan and Cunningham-Bussel, 2013). Under certain circumstances, ROS may be beneficial in promoting wound healing, and NADPH oxidase defects in patients with chronic granulomatous disease (Segal et al., 2011) and in animal models (Morgenstem et al., 1997) lead to aberrant inflammation. Chemical NADPH oxidase inhibitors such as apocynin have antiinflammatory effects in arthritis models (Hart et al., 1990) and reduce ethanol-induced liver inflammation in mice (Gustot et al., 2006). Disrupting the gp91phox-p47phox interaction using a peptide inhibits NADPH oxidase (Rey et al., 2001). Likewise, targeting PCNA could be a strategy to dampen this neutrophil contribution to inflammation. In a therapeutic setting, chemical targeting of NADPH oxidase protein interaction potentially allows temporary and dose-dependent reduction of ROS, favoring the dampening of inflammation compared with the complete loss of NADPH oxidase function. It should be noted that in the current study along with the antiinflammatory effect, short T2AA treatment not only did not interfere with epithelial cell proliferation in TNBS-induced colitis, but it significantly enhanced epithelium regeneration. Our data strongly suggest that T2AA in this setting primarily targeted inflammatory neutrophils and did not demonstrate a negative effect on regenerating tissue at the site of inflammation. Here we show that T2AA also reduced neutrophil survival as an added antiinflammatory mechanism to limit neutrophil-mediated tissue damage, as previously demonstrated for roscovitine, a cyclin-dependent kinase inhibitor (Rossi et al., 2006). Down-regulating neutrophil activation and promoting neutrophil apoptosis represent important mechanisms in inflammation resolution (Serhan et al., 2007; Jones et al., 2016). Nonetheless, evaluation of the therapeutic value of T2AA as an antiinflammatory drug will require investigation of its effects on other models of inflammation. Notably, T2AA also did not adversely affect neutrophil progenitor survival. As previously mentioned, chemical strategies are reversible, and moderate reduction of NADPH oxidase function may provide antiinflammatory effects without compromising antipathogen function.

Cytosolic PCNA acts as a molecular “organizer” of neutrophil effector functions, coordinating apoptosis and governing neutrophil activation. Neutrophils represent a unique cellular model to elucidate the functions of cytosolic PCNA, and these findings could lead to the concept of molecular linkage of cell survival, oxidant-related processes, and metabolism that is relevant to the control of inflammation.

Materials and methods
Isolation of human neutrophils
Neutrophils were obtained from EDTA-anticoagulated blood from healthy donors in agreement with Etablissement Francais du Sang guidelines. Each donor gave written informed consent to participate in this study, which was approved by the Institut National de la Santé et de la Recherche Médicale Institutional Review Board (Paris, France). The study was conducted according to the Declaration of Helsinki. Neutrophils were isolated in LPS-free conditions by dextran sedimentation and Ficoll centrifugation as previously described (Witko-Sarsat et al., 2010).

Identification of PCNA partners by MS
Control IgG or rabbit polyclonal anti-PCNA Ab6 antibody (20 μg) was cross-linked to 50 μl of protein G Sepharose beads (Pierce) using 100 μl dimethyl pimelimidate (20 mM; Pierce) in cross-linking buffer (200 mM carbonate buffer, pH 9) for 30 min according to the manufacturer’s instructions and as previously described (Borregaard et al., 1983; Ohayon et al., 2016). Proteins (200 μg) from human neutrophil cytosols were obtained by nitrogen cavitation to prevent granule rupture, preserving cytosolic proteins from artifactual proteolysis as previously described (Witko-Sarsat et al., 2010). Neutrophil cytosol was added to Sepharose beads cross-linked with anti-PCNA antibody and incubated for 2 h at 4°C. Proteins bound to the beads were recovered after extensive washing and were denatured at 95°C in reducing conditions with Laemmlisi buffer. Supernatants were loaded onto an SDS-PAGE gel for a short migration, and proteins were trypsin-digested in-gel. For liquid chromatography/MS protein analysis, peptides were concentrated, washed, and analyzed using a reverse-phase C18 column on a u3000 nanoHPLC connected to a Linear Trap Quadrupole-Orbitrap mass spectrometer (Thermo Fisher Scientific). LTQ MS/MS collision-induced dissociation spectra were acquired from the s20 most abundant ions detected in the Orbitrap MS scan. Protein identifications were performed with the Proteome discoverer 1.3 (Thermo Fisher Scientific) and with Mascot (http://www.matrixscience.com). Separate analysis was conducted and compared using MyPROMS software. The proteomic analysis was performed in the Proteomics 3P5 facility of the Cochin Institute–Paris Descartes University in Paris.

SPR
PCNA was expressed in E. coli and purified as previously described (Desplancq et al., 2016), except that size-exclusion chromatography was performed in 20 mM Tris, pH 7.5, 0.1 M NaCl, and 0.1 mM 2-ME. Recombinant human p47phox, p67phox, and Rac2 were produced as previously described (Raad et al., 2009). Recombinant p40phox expressed in E. coli was purchased from Creative Biomart. To generate the PX domain of p47phox, a truncated p47phox (residues 2-132) was produced in E. coli using the pGEX plasmid (generous gift of Dr. Yaffe, Massachusetts Institute of Technology, Cambridge, MA). SDS-PAGE analysis of purified recombinant proteins expressed in E. coli confirmed the purity and the expected molecular mass of each protein: PCNA (36 kD), p47phox (47 kD), p47phoxPX (14 kD), p67phox (67 kD), and Rac2 (22 kD). Interactions between PCNA and cytosolic proteins of the NADPH oxidase complex were studied by SPR on a Biacore T200 or a Biacore 3000 system (GE Healthcare) designed to calculate affinity and kinetic parameters. PCNA was covalently immobilized on a dextran layer...
sensor chip (CM5) with 10 mM Hepes and 150 mM NaCl, pH 7.4. The recombinant proteins p47phox, p47phox PX domain, p67phox, p40phox, and Rac2 at the indicated concentrations (25–400 nM) were used as analytes in a running buffer (50 mM Tris and 150 mM NaCl, pH 7.4) and were injected at 20 µl/min for 120 s over immobilized PCNA (4,100 response units [RU]) and over a control flow cell submitted to the coupling steps without PCNA. Binding experiments were performed at 25°C. In all experiments, the specific binding signal was obtained by subtracting the background signal.

For global fitting of the full-length p47phox
We used kinetic titration on the Biacore T200 system, referred to as single-cycle kinetics, to collect binding data for kinetic analysis. This method involves sequential injections of several analyte concentrations over the ligand immobilized on the sensor chip surface without a regeneration step between successive injections. The association (k_a) and dissociation (k_d) rate constants and K_D were calculated using BIASuite software.

For peptides and p47phox PX domain–PCNA interactions
The running buffer was PBS, pH 7.4 (peptides), or 50 mM Tris and 150 mM NaCl, pH 7.4 (p47phox PX), and the surface was regenerated with puls of 0.5 mM glycine, pH 3. The immobilization level for PCNA was 4,300 RU. Association phases ran for 120 s, and dissociation phases ran for 120 s on a Biacore 3000 system.

Inhibition assays
The phospholipid phosphatidylinositol (3,4,5)-trisphosphate (PIP3,4,5; Sigma-Aldrich), which specifically binds to the PX domain of p47phox, was used to inhibit the binding of p47phox to PCNA at the indicated concentrations ranging from 4 to 80 µM. The dose-dependent inhibition by the phospholipid PIP3,4,5 was performed on 4,100 RU of immobilized PCNA using the Biacore T200 apparatus. The solution containing p47phox (200 nM) was incubated for 10 min at 25°C with lipids at various concentrations before injection. The signal was measured at the end of the association phase (120 s) and compared with those obtained in the absence of lipids.

Inhibition studies using p21 peptide and p47phox peptides were performed using the Biacore 3000 on 2,600 RU of immobilized PCNA. p47phox (500 nM) samples were co-injected with peptides at various concentrations, and the association phase ran for 120 s. Because peptides interact with PCNA, we evaluated inhibition by measuring the SPR signal at the end of an extended dissociation phase (380 s), a time point at which peptides that have higher dissociation rate than the protein are completely dissociated and, consequently, at which the signal is due to p47phox only. Inhibition was then evaluated by comparing the remaining signal in the dissociation phase obtained when peptide and protein were co-injected to those measured after injection of the protein alone.

Kinetic values determination
Data were analyzed by global fitting to binding models of both the association and the dissociation phases for several concentrations simultaneously using BIASuite software. The simplest 1:1 Langmuir binding model was first tested. The two-state reaction model and heterogeneous ligand model were used when they significantly increased the statistical goodness-of-fit as calculated by BIASuite software. In the case of P47–PCNA interaction, the two-state reaction model suggested a possible conformational change of P47 during interaction. For short peptides that are supposed to be homogeneous in solution, the heterogeneous ligand model accounts for the presence of two sites that bind analyte independently of each other. SPR analysis was performed at the Structural Biology Institute at the Grenoble Instruct–European Research Infrastructure Consortium facility in Grenoble.

Crystallographic data collection and structure determination
Crystals were obtained using hanging drop vapor-diffusion technique at 20°C and by mixing 1 µl of PCNA at 4.7 mg/ml with 1 µl of the crystallization solution in the presence or absence of p47phox-106-127 peptide (Fig. S1). A search for PCNA crystallization conditions was based on previously published conditions and systematic screening using commercial screens. The best crystals were obtained with a crystallization solution containing 19–23% PEG 3350, 0.2–0.3 M NaCl, and 0.1 M Tris–HCl, pH 8.5. Approximately 12 h before crystal freezing, 100–500 µM of p47phox-106-127 peptide was added to the crystallization drop. 25% glycerol was added to the crystallization drop to prevent ice formation, and crystals were fished out and immediately frozen in liquid nitrogen. Crystallographic data were collected at ~160°C on beamlines BM30-A and ID30-A1 at the European Synchrotron Radiation Facility in Grenoble, France. Data processing was performed using the XDS package, and the PCNA structure was determined by the molecular replacement technique using Phaser and the CCP4 suite of programs. The search model used was the human PCNA (PDB entry 1AIX). The final PCNA model was obtained after several cycles of manual model building with COOT software and maximum likelihood refinement using the REFMACs program. The highest-resolution structure was refined against 2.8-Å data, but showed no significant electron density for the soaked peptide. The structure that showed the largest amount of (F_Obs − F_Calc) residual electron density, located in the vicinity of the interdomain-connecting loop, was refined at 3.22-Å resolution. No peptide could be reliably refined into this electron, and the final model accounts for only the trimeric PCNA structure. Data processing and refinement statistics are presented in Table S1.

Molecular modeling
Molecular modeling was used to propose two possible docking modes of the p47phox PX domain onto the PCNA. Models are based on the crystallographic structures of the human PCNA in complex with p21-141-160 peptide (PDB entry 1AXC) and of the human p47phox PX domain (PDB entry 1KQ6). For the first model, the 106–127 PX segment was manually modified with COOT software to have the PX 114–118 region adopting the same conformation as the 147–151 segment in p21 peptide, when bound to PCNA, with an orientation compatible with docking onto PCNA. The stereochemistry of the 106–127 PX segment was calculated using BIAevaluation software.

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optimized by the refinement procedure implemented in COOT. The second model was built by simply superposing the PX 122–126 helix turn onto the 147–151 segment in p21 peptide, when bound to PCNA.

**Transfection and differentiation of PLB985 cells**

PLB985 cell lines were cultured in RPMI supplemented with 10% FBS, 1-glutamine (2 mM), antibiotics (penicillin 100 U/ml and streptomycin at 100 µg/ml), and Hepes buffer. PLB985 cell granulocytic differentiation was induced by exposure to 0.5% DMF for 5 d and was validated by FACS analysis of CD11b expression, as described previously (De Chiara et al., 2013). For PCNA overexpression, PLB985 cells were stably transfected with empty pcDNA3 plasmid (pCT), pcDNA3-PCNA (PCNA), and pcDNA3-SV40NLS-PCNA (PCNA-NLS; Bouyad et al., 2012) to generate control PLB985, PLB985 overexpressing PCNA WT, and PLB985 overexpressing exclusive nuclear PCNA, respectively. For p21/waf1 overexpression, PLB985 cells were stably transfected with empty PRC-RSV-neo plasmid (Invitrogen) or PRC-RSV/p21A45R to generate control PLB985, PLB985SP21, and PLB985SP21A45R as previously described (Dublet et al., 2005). Notably, because p21/waf1 is sensitive to degradation by neutrophil proteinase 3, we have used the mutant p21A45R, which is resistant to serine proteases and is more stable in the current study (Martin et al., 2016). Western blot analysis using anti-PCNA (rabbit polyclonal Ab5) or p21/waf1 (rabbit polyclonal Santa Cruz) on PLB985 lysate was used to assess the expression of PCNA and p21/waf1, respectively. PCNA, p21/waf1, and actin bands were quantified and normalized as PCNA/β-actin ratio. Of note, the PCNA protein expression was not affected by the ectopic expression of p21/waf1.

For PCNA siRNA experiments, PLB985 cells were treated with DMF to induce granulocytic differentiation and transfected with 1 µM siRNA (Applied Biosystems) twice at a 24-h interval (day 4 and day 5 after DMF treatment) using the Amamax system (Lonza), according to the manufacturer’s instructions. In brief, 2 × 10^6 cells were resuspended in 100 µl of cell line solution with 1 µg of plasmid. The cells were then electroporated and transferred to culture plates. Transfected cells were cloned and selected based on their resistance to 1 mg/ml neomycin. FITC-conjugated control siRNA, used to detect intracellular siRNA by FACS and to monitor transfection efficiency, showed that the fluorescent siRNA was taken up in >90% of the PLB985 cells. Western blot analysis using anti-PCNA (rabbit polyclonal Ab5) on PLB985 lysate was used to validate the efficacy of PCNA siRNA in decreasing the expression of PCNA protein. PCNA and actin bands were quantified and normalized as PCNA/β-actin ratio. Flow cytometry analysis was performed at Cytometry and Immunobiology at the Cochin Institute in Paris.

**Measurement of NADPH oxidase activation using cell-free system experiments**

NADPH oxidase activity was measured in a cell-free system using membrane and cytosolic fractions of resting neutrophils as previously described (Molshanski-Mor et al., 2007). The reaction mixture contained 1 mM ATP, 1.25 mM EGTA, 50 µM guanosine 5′-(γ-thio)triphosphate (Sigma-Aldrich), and 7 × 10^6 cell membrane equivalent. When indicated, the p21 peptide or T2AA was preincubated for 10 min at 37°C at the indicated concentration. Cells were then stimulated with (S) or without (R) 100 µM Li-SDS (Fluka) for 5 min at 37°C before starting the reaction. The reaction was then initiated with 200 µM β-NADPH (Sigma-Aldrich) and cytochrome c (1 mg/ml). Absorbance at 550 nm was read immediately, and superoxide anion production was calculated using an extinction coefficient of 21.1 mM^−1 cm^−1 for reduced ferricytochrome c. The specificity of the reaction was evaluated by adding 20 µM DPI or 30 U/ml SOD. Results are expressed as nanomoles of O_2^−/min/7 × 10^6 cell membrane equivalent.

**Translocation of p47phox from the cytosol to the neutrophil membrane**

After isolation from whole blood, 100 × 10^6 neutrophils were resuspended in 5 ml of HBSS/+/− and incubated with or without T2AA (25 or 50 µM) at 37°C for 1 h. Neutrophils were then stimulated with PMA (200 ng/ml) for 8 min while the mixture was gently shaken. Ice-cold buffer was added, and cells were pelleted by centrifugation at 400 g for 8 min at 4°C. The pellets were resuspended in relaxation buffer containing 1 mM ATP, 1 mM EGTA, 0.5 mM PMSF, 25 nM Calyculin, and 5 µg/ml of aprotinin, leupeptin, and pepstatin. The cells were then disrupted by sonication (twice for 10 s each time) at 4°C and centrifuged at 400 g for 8 min. The postnuclear supernatant was loaded onto a discontinuous sucrose gradient (35 and 15% sucrose) diluted in relaxation buffer and centrifuged for 45 min at 150,000 g. The cytosolic and membrane fractions (in the upper and the intermediate layers, respectively) were collected. Proteins were analyzed by Western blot analysis using
neutrophils spotted on poly-L-lysine coated glass slides and allowed to dry in a humid chamber by gravity sedimentation, then fixed with 2% paraformaldehyde for 10 min, permeabilized with 0.2% Tween 20 for 15 min at 37°C, and blocked with 5% BSA in PBS. Cells were then incubated overnight at 4°C with a rabbit anti-PCNA polyclonal antibody (1:200) and mouse anti-p47phox mAb (1:200) diluted in 1% BSA/PBS. After washing, cells were incubated with Alexa Fluor 488-conjugated (green) goat anti-rabbit antibody (1:200) and Alexa Fluor 568-conjugated (red) goat anti-mouse antibody (1:200) for 1 h at room temperature in the dark. Nuclei were stained with TO-PRO-3 iodide (Invitrogen). A proximity ligation assay using the Duolink system was performed on human isolated neutrophils spotted on poly-L-lysine-coated glass slides according to the manufacturer’s instructions using two primary antibodies from different species (the rabbit polyclonal anti-PCNA Ab5, the mAb anti-p47phox [Millipore], or the rabbit polyclonal anti-rac 2 [Abcam]). When indicated, neutrophils were treated with PMA (0.1 μg/ml for 10 min) before labeling. Resting neutrophils treated with anti-p47phox; anti-PCNA antibodies show a cytosolic fluorescence indicative of a colocalization. Hoechst was used for nuclear staining. Stained cells were examined with a Zeiss LSM510 confocal microscope (63×/1.4 numerical aperture objective), and the images were imported into an LSM Image browser for analysis. The designation “merge” for indirect immunofluorescence analysis corresponds to colocalization of PCNA and p47phox. The magnification used was 63×, and different zoom settings were used. The final magnification is indicated by the presence of a scale bar for each experiment.

**Coimmunoprecipitation experiments**

Human isolated neutrophils (25 × 10⁶/ml per condition) were stimulated with PMA (0.1 μg/ml) or f-MLF (1 μM) for 1 h at 37°C with 5% CO₂. Neutrophil cytosol was obtained by sonication in a hypotonic Hepes buffer (50 mM) supplemented with inhibitors (4 mM PMSF, 20 mM leupeptin, 20 mM pepstatin, 270 mM orthovanadate, 200 mM EGTA, and 500 mM EDTA) with a Soniprep 150 Plus sonicator (1 Hz for 10 s). Protein concentration was measured using the BCA kit (Pierce). Coimmunoprecipitation experiments were performed on cytosol (500 μg of protein) by adding 7.5 μl rabbit anti-PCNA Ab5 or control rabbit IgG for 15 min at 4°C under rolling. Next, 50 μl of protein A beads were added and incubated for 30 min at 4°C under rolling and 15 min on ice. The sample was loaded onto a column containing Sepharose-coated magnetic beads (Miltenyi) prehydrated with a washing buffer (50 mM Tris, pH 8, 300 mM NaCl, and 1% NP-40). The column was washed four times with this salt-containing buffer and five times with a wash buffer (20 mM Tris, pH 7.5). The sample was eluted from the column with 50 μl 5x boiling sample buffer. Cytosol (30 μg of protein) elution and exclusion samples were analyzed by Western blotting using anti-p47phox (BD Biosciences; 610354) and PCNA (PC10 mAb) followed by HRP-conjugated anti-mouse IgG (Nordic Immunology; diluted 1:5,000), using the SuperSignal West Pico detection kit (Pierce) as previously described (Witko-Sarsat et al., 2010).

**Mouse models of inflammation**

Animal studies were performed in accordance with the European Community Guidelines (authorization number B-75-1394 and C-75-1801). All protocols were approved by Ethics Committee for Animal Research of Université Paris Descartes (CEEA34-VWS.074.12) and of Faculté Bichat (CEEA-JCM.121).

**Zymosan-induced peritonitis in mice**

The model of zymosan-induced peritonitis was used to examine the NADPH oxidase activity in murine inflammatory neutrophils. Peritonitis was induced in mice (male C57BL/6J mice, 9–30 wk old; Charles River) by an intraperitoneal injection of 1 mg of Zymosan A from Saccharomyces cerevisiae (40 mg/kg; Sigma-Aldrich) in 0.5 ml of sterile PBS as previously described (Martin et al., 2016). After 4 h, mice were sacrificed by cervical dislocation, and the peritoneal cavities were washed with 4 ml of PBS. Aliquots of lavage fluids (10 μl) were diluted in Turk’s solution (0.01% crystal violet in 3% acetic acid), and cells were counted using a Malassez hemocytometer. Analysis of neutrophils or macrophages was performed by flow cytometry after immunolabeling with anti-Ly6G-FTTC (BD Biosciences) and anti-F4/80–Alexa Fluor 647 (AbD Serotec), respectively. Isolated neutrophils (1 × 10⁶) from the peritoneal lavage were suspended in 0.5 ml HBSS+/+ and stimulated with or without PMA (1 μg/ml), and luminol CL was recorded by a TriStar LB 941 Multimode Microplate Reader as described above. When indicated, zymosan-induced peritonitis was performed in p21/Waf1 knockout mice (p21⁻/⁻) obtained from the Jackson Laboratory (B6;129S2-Cdkn1atm1Tyj/J) and housed with their WT littermate controls (Martin et al., 2016). To evaluate the effect of in vivo injection of T2AA on oxidative burst in inflammatory neutrophils, mice were pretreated with or without T2AA (20 mg/kg) for 30 min and 17 h before the injection of zymosan to trigger peritonitis.

To evaluate the ex vivo effect of T2AA on the survival of neutrophils in C57BL/6J WT mice compared with mice lacking p47phox (p47phox⁻/⁻ mice [Ncf1⁻/⁻]); the Jackson Laboratory), peritonitis was induced in mice (male and female C57BL/6J mice, 9–30 wk old) by an intraperitoneal injection of zymosan. After 4 h, mice were sacrificed with isoflurane, and the peritoneal cavities were washed with 4 ml of PBS. Neutrophils in C57BL/6J WT mice compared with mice lacking p21/Waf1 (p21⁻/⁻) obtained from the Jackson Laboratory (B6;129S2-Cdkn1atm1Tyj/J) and housed with their WT littermate controls (Martin et al., 2016). To evaluate the effect of in vivo injection of T2AA on oxidative burst in inflammatory neutrophils, mice were pretreated with or without T2AA (20 mg/kg) for 30 min and 17 h before the injection of zymosan to trigger peritonitis.

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Induction of TNBS-induced colitis in mice
We investigated the therapeutic effect of T2AA in TNBS-induced colitis in mice as previously described (Krimi et al., 2008). Male C57BL/6J mice (19–22 g; Janvier Laboratories) were deprived of food for 24 h before colitis induction. Then mice were anesthetized for 90–120 min for the induction of colitis. They received an intrarectal administration of TNBS (40 µl, 150 mg/kg) dissolved in a 1:1 mixture of 0.9% NaCl with 100% ethanol. Control mice received a 1:1 mixture of 0.9% NaCl with 100% ethanol. After each rectal administration of TNBS, mice were kept in an inverted position for 30 s. 6 and 24 h after TNBS injection, mice received an intraperitoneal injection of T2AA (25 mg/kg) or vehicle solution. Animals were sacrificed 2 d after TNBS administration. When indicated, neutrophil depletion was performed using two injections of anti-Ly6G antibody (clone 1A8) as previously described (Coffelt et al., 2015), 18 h( 400 µg/mouse) before and 5 h(100 µg/mouse) after induction of colitis (Fig. 8A).

Colonic inflammation was evaluated by the weight and length of each intestine (anus to cecum). The colons were opened and examined to evaluate macroscopic lesions according to the Wallace criteria (Wallace et al., 1989). The Wallace score rates macroscopic lesions on a scale from 0 to 10 based on features reflecting inflammation such as hyperemia, thickening of the bowel, and extent of ulceration. For histological analysis, colon specimens located 1 cm above the anus were removed to be included in paraffin. Several 5-µm paraffin sections were prepared and stained with H&E according to the Ameho criteria (Ameho et al., 1997). This grading is scaled from 0 to 6 and takes into account the presence of erosion, ulceration, or necrosis and the depth and surface of lesions. Immunohistological labeling of colon sections was also performed with a polyclonal rabbit anti-MPO (A0398, Dako) and with the neutrophil-specific marker antibody (rat mAb clone 6A608; Santa Cruz) as previously described (Martin et al., 2016). To evaluate the presence of neutrophils within the colon, we quantified the immunostaining obtained with the neutrophil marker (clone 6A608). Slides were entirely scanned by the Lamina Slide Scanner (PerkinElmer), and regions of interest were manually defined using Case Viewer (https://www.3dhistech.com/software-downloads/). Surfaces of tissue regions of interest were determined with INFORM software (PerkinElmer) by tissue segmentation. Then, surfaces of neutrophil signal were determined by the “threshold” command. The ratio between these two surfaces were calculated to analyze and compare the variations of the immunostained surfaces percentage. The immunohistochemistry study was performed at the Histology, Immunostaining, and Laser Microdissection facility at Cochin Institute in Paris.

Noninvasive in vivo imaging of NADPH oxidase activity using L012-enhanced CL
Measurement of in vivo neutrophil NADPH activation was performed as previously described (Kielland et al., 2009), with some modifications. For the zymosan-induced peritonitis model, mice were first injected intraperitoneally with 1 mg zymosan to induce peritonitis, followed by T2AA treatment (25 mg/kg) 4 h later. To measure in vivo neutrophil NADPH activation, the luminescent probe L012 (Sobioda) was prepared freshly right before the experiment by dissolving 2 mg L012/ml in sterile PBS and kept in the dark. L012 was administrated at 20 mg/kg by intraperitoneal injection to the right flank of isoflurane-anesthetized (1.5–2.5%) mice 6 h after the initial injection of zymosan. In the model of TNBS-induced colitis, L012 was injected at 24 and 48 h after TNBS treatment to measure in vivo activity of NADPH oxidase at these time points. Results are expressed as the variation of L012-CL between 24 and 48 h. For all the in vivo imaging of NADPH oxidase activity, L012 was administered to mice that had been under anesthesia for 3 to 4 min, after which mice were placed into the light chamber of the Photon Imager RT imaging system (BiospaceLab) equipped with an intensified charge-coupled device camera that allows real-time recording of emitted photons. The number of photons produced by L012 during in vivo imaging was collected during 10 min. A maximum of five individual mice per acquisition were positioned supine in the chamber. Regions of interest were drawn using M3 Vision proprietary software (BiospaceLab). This software allows quantification of photons emitted over time in each region of interest with real-time photon collection. Quantification was performed measuring emitted photons in a time window corresponding to the plateau of the intensity curve, for each region of interest. Light emission from the abdominal region of interest was quantified and expressed as photons/s/cm²/steradian. This study was performed at the Small Animal Imaging facility at the Cochin Institute in Paris.

Statistical analysis
Statistical analysis was performed using the Statview software package. Comparisons were made using the Student’s t test, multicomparison ANOVA, or the Mann-Whitney U test as indicated. Differences were considered significant at P < 0.05.

Accession numbers
The structure factors and coordinates have been deposited in the PDB under accession nos. 6FCM (PCNA structure at 2.8 Å) and 6FCN (PCNA structure at 3.22 Å in combination with the p47phox peptide).

Online supplemental material
Fig. S1 is a structural analysis of the association between PCNA and the p47phox peptides. Fig. S2 shows the effect of PCNA inhibition on viability in differentiated PLB985 cells or human neutrophils. Fig. S3 shows the effect of T2AA on degranulation, chemotaxis, and phagocytosis of E. coli in isolated human neutrophils. Fig. S4 shows the effect of T2AA on neutrophil survival isolated from WT and from p47phox−/− mice. Table S1 shows crystallographic data collection and refinement statistics.

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