Identification of a Ligand-induced Transient Refractory Period in Nuclear Factor-κB Signaling*

Britney L. Moss‡1,2, Shimon Gross‡1, Seth T. Gammon3, Anant Vinjamoori4, and David Piwnica-Worms‡5,6

From the ‡1Molecular Imaging Center, Mallinckrodt Institute of Radiology, and the ‡5Department of Molecular Biology and Pharmacology, Washington University School of Medicine, St. Louis, Missouri 63110

In response to a variety of extracellular ligands, nuclear factor-κB (NF-κB) signaling regulates inflammation, cell proliferation, and apoptosis. It is likely that cells are not continuously exposed to stimulating ligands in vivo but rather experience transient pulses. To study the temporal regulation of NF-κB and its major regulator, inhibitor of NF-κBα (IκBα), in real time, we utilized a novel transcriptionally coupled IκBα-firefly luciferase fusion reporter and characterized the dynamics and responsiveness of IκBα processing upon a short 30-s pulse of tumor necrosis factor α (TNFα) or a continuous challenge of TNFα following a 30-s preconditioning pulse. Strikingly, a 30-s pulse of TNFα robustly activated inhibitor of NF-κB kinase (IKK), leading to IκBα degradation, NF-κB nuclear translocation, and strong transcriptional up-regulation of IκBα. Furthermore, we identified a transient refractory period (lasting up to 120 min) following preconditioning, during which the cells were not able to fully degrade IκBα upon a second TNFα challenge. Kinase assays of IKK activity revealed that regulation of IKK activity correlated in part with this transient refractory period. In contrast, experiments involving sequential exposure to TNFα and interleukin-1β indicated that receptor dynamics could not explain this phenomenon. Utilizing a well accepted computational model of NF-κB dynamics, we further identified an additional layer of regulation, downstream of IKK, that may govern the temporal capacity of cells to respond to a second proinflammatory insult. Overall, the data suggested that nuclear export of NF-κB/IκBα complexes represented another rate-limiting step that may impact this refractory period, thereby providing an additional regulatory mechanism.

Adequate resolution of an inflammatory reaction is as equally important as initiation. Persistent or fulminant responses can cause detrimental consequences both locally and systemically (1), and resolution of inflammation is important for both termination of an acute response as well as for prevention of destructive chronic responses. It is therefore not surprising that mechanisms aimed at rapid and specific initiation of proinflammatory reactions have co-evolved with mechanisms that provide timely termination of such processes. From a systems biology perspective, such “switchability” can be achieved by intracellular feedback loops that permit ligand-induced desensitization and resensitization of proinflammatory signaling cascades (2).

In this regard, recent studies have shown that nuclear factor-κB (NF-κB) signaling plays a critical role in both initiation and resolution of inflammation (2, 3). The transcription factor NF-κB is a key regulator of innate and adaptive immune responses as well as a mediator of cell survival and proliferation (4). Improper regulation of NF-κB contributes to induction and progression of a wide range of human disorders, including a variety of pathological inflammatory conditions, neurodegenerative diseases as well as many types of cancer (5, 6). In resting cells, inactive NF-κB is sequestered in the cytoplasm by binding to members of the inhibitor of NF-κB (IκB) family. Canonical activation of NF-κB depends on IκB kinase (IKK)-regulated proteasomal degradation of IκBα, an event that frees NF-κB for nuclear translocation within minutes (4, 7). Upon nuclear transport, NF-κB regulates the transcription of a few hundred genes (8–10) that can be divided into four major families (10, 11): 1) proinflammatory genes (e.g. cyclooxygenase 2, interleukin-1 (IL-1), tumor necrosis factor α (TNFα), inducible nitric-oxide synthase, intercellular adhesion molecule-1, E-selectin, etc.), 2) proproliferative genes (e.g. cyclin D, and cellular myelocytomatosis), 3) antiapoptotic genes (B-cell leukemia/lymphoma 2, B-cell leukemia/lymphoma extra long, X-linked inhibitors of apoptosis protein, and cellular inhibitors of apoptosis protein), and 4) autoinhibitory genes (e.g. A20, cylindromatosis, suppressor of cytokine signaling 1, and IκBα).

With respect to the last, other transcriptionally independent processes, aimed at autoinhibition of NF-κB activity, do exist. Such mechanisms down-regulate NF-κB signaling on a much shorter time frame (seconds to minutes). These include homologous receptor desensitization (12, 13), asymmetric heterologous receptor desensitization (13, 14), autocatalytic C-terminal IKK hyperphosphorylation (15), and protein phosphatase 2C-dependent dephosphorylation of IKK (16).

* This work was supported in part by National Institutes of Health Molecular Imaging Center Grant P50 CA94056. The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked “advertisement” in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

1 Both authors contributed equally to this work.

2 Supported in part by a National Science Foundation graduate research fellowship grant.

3 To whom correspondence should be addressed: Mallinckrodt Institute of Radiology, Washington University School of Medicine, 510 S. Kingshighway Blvd., Box 8225, St. Louis, MO 63110. Tel.: 314-362-9359; Fax: 314-362-0152; E-mail: piwnica-wormsd@mir.wustl.edu.

4 The abbreviations used are: NF-κB, nuclear factor-κB; IκBα, inhibitor of NF-κBα; IKK, IκB kinase; IL-1, interleukin-1; TNFα, tumor necrosis factor α; FLuc, firefly luciferase; GST, glutathione S-transferase; PBS, phosphate-buffered saline; DME, Dulbecco/Vogt modified Eagle’s minimal essential medium; IKK-KA, IKK kinase assay; C, continuous; P, pulse; P + C, pulse followed by continuous; EGFP, enhanced green fluorescent protein.
Refractory Period in NF-κB Signaling

Considering the complex nature of the inflammatory milieu, one would expect that stationary tissue-residing cells are exposed to a myriad of temporally distinct NF-κB-stimulating cues. For instance, cells can be directly stimulated by pathogen-derived products (e.g. lipopolysaccharide through TLR4 (toll-like receptor 4) receptors (17)), exposed to numerous soluble proinflammatory stimuli produced by circulating effector cells (e.g., cytokines, chemokines, etc.), and/or experience inflammation-induced oxidative stress (18). These signals can occur simultaneously or sequentially to one another. For example, systemic administration of bacterial lipopolysaccharide to mice was shown to induce transient production of TNFα (serum levels peaking at ~1.5 h and quickly returning to base line), but IL-1β production was delayed and prolonged (first detected at 2 h, but lasting >5–6 h) (19). Thus, cells co-expressing TLR4, IL-1, and TNFα receptors would sequentially interrogate signals arising from lipopolysaccharide, TNFα, and IL-1β, each of which could independently activate NF-κB.

Central to any signaling desensitization mechanism is a refractory period during which cells cannot fully respond to a second insult (autologous or heterologous desensitization). Therefore, consideration of the dynamic pattern of stimulus exposure described above begs the immediate question of whether cells can instantly initiate an NF-κB response to a second activating stimulus, and if not, when will such cells be able to remount a full response again? Specifically, are ligand-preconditioned cells capable of eliciting NF-κB activation to the same extent as naïve cells?

Little is known about the capacity of cells to activate NF-κB in response to a second activating challenge, since the highly dynamic nature of this process presents many technical difficulties. These include low temporal resolution of conventional transcriptionally dependent NF-κB reporter gene assays, low throughput, inability to acquire longitudinal data, and the semi-quantitative nature of traditional biochemical assays (e.g. electrophoretic mobility shift assay, immunoblotting, etc.). Such limitations render these assays incapable of accurate analysis of the early, ligand-induced dynamic changes in the capacity of cells to elicit a response to a second challenge.

To efficiently address this question, we generated an improved, transcriptionally coupled version of a previously published genetically encoded IκBα-firefly luciferase (IκBα-FLuc) fusion reporter (20) in conjunction with dynamic, live cell bioluminescence imaging of cultured cells. We chose to focus on HepG2 human hepatoma cells as a model system, because 1) NF-κB signaling has been extensively studied in these cells, 2) HepG2 cells have been shown to activate NF-κB in response to a variety of proinflammatory ligands (21), 3) these cells can be easily transfected with readily available reagents, and, most importantly, 4) the pivotal role that NF-κB signaling plays in hepatocytes to regulate inflammation, apoptosis, and carcinogenesis (22).

Using bioluminescence imaging of live cells in conjunction with a variety of biochemical assays, we demonstrate herein that a 30-s preconditioning exposure to TNFα is sufficient to robustly activate IKK, culminating in IκBα degradation, NF-κB nuclear translocation, and strong transcriptional up-regulation of IκBα. Furthermore, the capacity of preconditioned cells to degrade IκBα in response to a second TNFα challenge is transiently refractory, regaining full responsiveness ~120 min later. Finally, both IKK regulation and possibly NF-κB nuclear export, but not receptor dynamics, govern this transient refractory period. This study highlights the interlocking layers of NF-κB regulation, ensuring efficient and timely propagation as well as termination of proinflammatory signals.

EXPERIMENTAL PROCEDURES

Materials—d-Luciferin (potassium salt) was from Biosynth (Naperville, IL). Human TNFα and IL-1β were from R&D Systems (Minneapolis, MN). Complete protease inhibitor mixture was from Roche Applied Science (Basel, Switzerland). [γ-32P]ATP was from PerkinElmer Life Sciences. Carbencillin, isopropyl β-D-1-thiogalactopyranoside, ampicillin, kanamycin, glutathione S-transferase (GST), β-glycerolphosphate, NaCl, NaF, Na3VO4, KOH, MgCl2, EDTA, phenylmethylsulfonyl fluoride, Nonidet P-40, Tween 20, Triton X-100, ATP, dithiothreitol, paraformaldehyde, cycloheximide, and HEPES were from Sigma.

Plasmids—pκB5→FLuc (Stratagene, La Jolla, CA) contains five repeats of a κB motif upstream of a minimal TATA box controlling expression of firefly luciferase. pκB5→IκBα-FLuc was produced by cloning an EcoRI/HpAl (blunt) fragment from pCMV→IκBα-FLuc (20) into the EcoRI and EcoRV (blunt) sites of pκB5→FLuc. pκB5→FLuc, pCMV→IκBα-FLuc, and pκB5→IκBα-FLuc were propagated in TOP10 electrocompetent Escherichia coli (Invitrogen) and purified using Qiagen HiSpeed Maxi Kits (Qiagen, Valencia, CA). pGST-IκBαN (encoding for GST fused to the N-terminal fragment of human IκBα (1–54)) was a kind gift from Prof. Alexander Hoffmann (University of California San Diego). pGST-IκBαN was propagated in BL21 codon + E. coli cells (Stratagene).

Cells and Transfections—HepG2 human hepatoma cells were from the American Type Culture Collection (Manassas, VA). Cells were cultured in DMEM supplemented with heat-inactivated fetal bovine serum (10%) and L-glutamine (2 mM). Cells cultures were grown at 37 °C in a humidified atmosphere of 5% CO2. HepG2 cells (104) were transiently transfected (Fugene 6; Roche Applied Science) with pκB5→IκBα-FLuc (200 ng/well) and plated in black-coated 24-well plates (In Vitro Systems GmbH, Gottingen, Germany). Cells were then allowed to recover for 48 h prior to imaging.

Dynamic Bioluminescence Live Cell Imaging—Prior to imaging, cells were washed with prewarmed phosphate-buffered saline (PBS, pH 7.4) and placed into 900 μl of colorless HEPES-buffered DMEM, supplemented as above and with d-luciferin (150 μg/ml). Cells were allowed to equilibrate for 1 h (37 °C) before proceeding with ligand stimulation and imaging. Four different stimulation regimens were included in this study. 1) For continuous TNFα (C), TNFα (final concentration 20 ng/ml) or vehicle (colorless DMEM) was added (100 μl) to d-luciferin-containing DMEM, and imaging was performed before and at the indicated time points after the addition of TNFα. 2) For TNFα pulse (30 s, P), cells were pulsed for 30 s with TNFα (20 ng/ml) or vehicle, washed with PBS, returned to d-luciferin-containing DMEM, and imaged before and at the indicated time points after the pulse of TNFα. 3) For TNFα preconditioning...
ing (30-s pulse) followed by continuous TNFα challenge (P + C), at t₀, cells were pulsed for 30 s with TNFα (20 ng/ml) or vehicle, washed with PBS, returned to d-luciferin-containing DMEM (900 μl), and imaged before and at the indicated time points after the pulse of TNFα. At t₀, TNFα (final concentration 20 ng/ml) or vehicle (colorless DMEM) were again added (100 μl), and imaging was performed before and at the indicated time points after the addition of TNFα. 4) TNFα preconditioning (30-s pulse) followed by continuous IL-1β challenge (P + C) was as in method 3, but continuous challenge was performed with IL-1β (10 ng/ml).

TNFα or IL-1β challenge was performed at the following challenge (C). Following this preconditioning (P). At t₀ + 25 min, time of maximal IkBα degradation (20); see Fig. 2A for schematic timeline), cells were harvested (by scraping) in reporter lysis buffer (Promega, Madison, WI). Cell lysates were normalized for protein content by a BCA protein assay (Promega, Madison, WI). Lysates were mixed with luciferase assay buffer (190 μl; 25 mM HEPES, 154 mM NaCl, 5.4 mM MgSO₄, 10 mM dithiothreitol, 5 mM ATP, 150 μg/ml d-luciferin, pH 8.0) in a 96-well plate immediately prior to imaging. Lysates were transferred to a scintillation vial, and radioactivity was determined on a β counter (Beckman Coulter, Fullerton, CA).

Calculating Ligand-dependent IKK Responsiveness—IKK responsiveness profiles (i.e. the net kinase capacity of IKK in response to a second challenge of TNFα, as a function of time after initial 30-s preconditioning) were calculated numerically from IKK-KA data using the following formula,

\[ \text{IKK responsiveness} = \frac{PC_{x+10} - P_{x+10}}{PC_0 \left( \frac{C_0}{C_{10}} \right)} \]

where \( PC_{x+10} \) is IKK activity of preconditioned plus challenged cells, as recorded 10 min postchallenge. \( P_{x+10} \) is the residual IKK activity of preconditioned but unchallenged cells.
at this exact time point. $C_0$ and $PC_0$ are initial IKK activities of challenged but unpreconditioned and fully preconditioned and challenged cells, respectively. $C_{10}$ is the maximal IKK activity of challenged but unpreconditioned cells (recorded 10 min post-challenge). Note that although all parameter units in the nominator and denominator are in counts/min, IKK responsiveness is dimensionless, similar to IkBα responsiveness.

Computational Simulations—To simulate the dynamics of major regulators on the IKK-NF-κB axis, we used a well-established computational model generated by Hoffmann et al. (25) and refined by Werner et al. (23). Briefly, an experimentally or hypothetically derived IKK activity profile was fed into the program as an input. Embedded in the model were 24 components, 70 reactions, and 70 parameters or rate constants for these reactions. Differential equations were solved numerically using Matlab 7.0 (Mathworks, Natick, MA) with subroutine Ode15s. Interpolated and extrapolated (0–360 min at 5-min intervals) IKK activity profiles were calculated (Origin version 7.5, OriginLab, Northampton, MA) from experimental IKK-KA data (see above). To fit the model, initial steady-state IKK activity (i.e. intracellular concentration of active IKK) was set to be 1 nm. To computationally simulate IkBα dynamics of cells challenged at different times after initial preconditioning, when assuming no upstream IKK or receptor regulation, we used hypothetical IKK activity profiles as inputs, derived from superimposing experimentally acquired IKK activity profiles of 30-s pulsed and continuously treated cells at increasing intervals (30, 60, 120, and 240 min; see Fig. 4, black lines).

Immunofluorescence Microscopy—HepG2 cells were seeded into 35-mm glass bottom culture dishes (MatTek Corp., Ashland, MA) and grown to ~40% confluence. Cells were pulsed for 30 s with TNFα as above and fixed at the indicated time points (by washing once with PBS, followed by fixation (4% paraformaldehyde for at least 15 min) and permeabilization (ice-cold methanol, 10 min at ~20 °C)). Cells were washed in PBS, blocked in 5% normal goat serum in 0.3% Triton X-100, PBS (1 h), and then incubated with anti-p65 antibody (Santa Cruz Biotechnology; 1:200 in 0.3% Triton X-100, PBS at 4 °C, PBS (1 h), and then incubated with anti-p65 antibody (Santa Cruz Biotechnology; 1:200 in 0.3% Triton X-100, PBS at 4 °C, overnight with rocking). Cells were next incubated with Alexa Fluor 635-conjugated goat anti-rabbit antibody (Invitrogen; 1:200 in 0.3% Triton X-100, PBS, 90 min, at room temperature with rocking). Cells were washed three times with PBS before being mounted with VECTASHIELD Mounting medium (Vector Laboratories; Burlingame, CA). Confocal images were captured using the ×40 objective (water immersion) on a Zeiss Axiovert 200 (Zeiss, Thornwood, NY) laser-scanning microscope equipped with the appropriate filter sets and analyzed using a Zeiss LSM Image Browser and Adobe Photoshop CS2.

RESULTS

Real Time Bioluminescence Imaging of pκB$_5$→IκBα-FLuc-expressing Cells Recapitulated IKK-induced Dynamics of Endogenous IκBα—To monitor ligand-induced IκBα rapid dynamics as well as physiologic transcriptionally coupled behavior, we modified our previous IκBα-FLuc fusion reporter (20) to be driven by a synthetic promoter composed of five tandem κB response elements (TGGGGACTTCCGC) followed by a minimal TATA-box. We hypothesized that this reporter would allow quantitative measurements of IKK-induced degradation as well as NF-κB-induced resynthesis and post-translational stabilization of IκBα from intact living cells (Fig. 1A). To validate use of this reporter, HepG2 cells were transiently transfected with a plasmid encoding the reporter and allowed to recover for 2 days before stimulation with a continuous or 30-s pulse of TNFα (20 ng/ml) to induce IKK activation. Upon the addition of TNFα, a rapid and dramatic decrease in bioluminescence was observed when readouts were normalized to untreated controls (20) under both continuous (C) and 30-s pulse (P) regimens (Fig. 1, B and C). This decrease in normalized bioluminescence, reflecting IKK-induced reporter degradation was followed by a sharp increase in bioluminescence, reflecting NF-κB-dependent reporter resynthesis, reaching maximum values at ~120 min and then gradually declining toward base line. Note that the rate at which IκBα levels return to base line is steeper under continuous TNFα treatment compared with the 30-s pulse, providing evidence for reactivation of ligand-induced IκBα degradation during continuous stimulation (23). The magnitude of the initial decrease in bioluminescence was greater in continuously treated cells than in 30-s pulsed cells (70% versus 40% of initial decrease, respectively), indicating that a 30-s pulse of TNFα leads to ~50% depletion of the IκBα-NF-κB pool compared with continuous TNFα exposure (Fig. 1C, 120 min). These data suggested that 1) this reporter construct could report on both IKK-induced IκBα degradation and successive resynthesis of IκBα, 2) a 30-s pulse of TNFα at a saturating concentration (20 mg/ml) elicited robust IKK activity, culminating in IκBα degradation and full IκBα transcriptional up-regulation, and 3) with the current κB$_5$ synthetic promoter system, there was a nonlinear relationship between IκBα degradation and NF-κB-dependent resynthesis of IκBα (i.e. saturation of IκBα resynthesis even at submaximal IκBα degradation levels).

Strikingly, Western blot analysis revealed that endogenous IκBα behaved exactly as the reporter under both C and P conditions, recapitulating the degree of degradation, recovery, and return to base line (Fig. 1D). Pretreating pκB$_5$→IκBα-FLuc-expressing HepG2 cells with cycloheximide did not affect degradation of IκBα-FLuc but abolished signal recovery, indicating that this phase was totally dependent upon transcription and translation of new IκBα-FLuc (Fig. 1E).

TNFα Preconditioning Induces a Transient Refractory Period of IκBα Processing—Upon a proinflammatory insult in vivo, effector cells (e.g. circulating macrophages) release TNFα and other activating cytokines in a temporally and spatially discrete manner. As a consequence, stationary target cells (e.g. epithelial cells, endothelial cells, hepatocytes, etc.) will sense a stochastic rise in the levels of such proinflammatory ligands. In such a dynamic environment, as ligand-secreting cells continuously migrate to sites of inflammation, it is anticipated that over time, target cells will experience multiple pulses of activating ligands.

We therefore aimed to elucidate the effects of such ligand pulses on the capacity of hepatocytes to respond to a subsequent challenge of the same ligand. Having shown that 1) pκB$_5$→IκBα-FLuc provided an accurate readout of IκBα processing in intact cells and that 2) a 30-s pulse was sufficient to induce robust IKK activity, we next sought to investigate...
whether a short 30-s preconditioning pulse with TNFα had a substantial effect on the capacity of cells to process IkBα upon a subsequent continuous TNFα challenge.

HepG2 cells transiently expressing pxB5→IkBα-FLuc were given a 30-s pulse of TNFα (20 ng/ml) or vehicle at t₀, washed, replaced in media containing D-luciferin and repeatedly imaged (every 5 min) prior to a TNFα challenge. At t₃₀₀, t₄₂₀₀ or t₃₄₀₀ (min) after pulsing, cells were then challenged with a second continuous concentration of TNFα (20 ng/ml), and live cell imaging was continued up to 360 min. To compare the processing dynamics of IkBα-FLuc in naive (un preconditioned) cells with that of preconditioned cells, the resulting bioluminescence profiles of preconditioned cells (Fig. 2A, black lines) were plotted along with the bioluminescence profile of unpreconditioned cells (i.e. only treated with continuous TNFα at t₀, red line, Fig. 2A). The different graphs represent the differential dynamics of IkBα-FLuc processing as the preconditioning pulse-challenge (P-C) intervals temporally increased (0–240 min).

We observed that challenging preconditioned cells with a continuous exposure to TNFα near the time that they had achieved maximal degradation from the preconditioning pulse (i.e. 30 min postpreconditioning) resulted in a small amount of additional IkBα degradation. As the interval between preconditioning and challenge increased, the magnitude of challenge-induced IkBα degradation also increased. These data suggested that the TNFα-NF-κB system possessed a built-in refractory period following TNFα treatment that prevented cells from fully responding to a second exposure to ligand. To quantify this phenomenon independent of confounding factors that may affect dynamic bioluminescence readouts (e.g. D-luciferin, ATP, O₂, or pH dynamics) and to verify its existence for endogenous IkBα, we performed a similar experiment, but instead of live cell imaging, we harvested whole cell lysates at tₓ + 25 min (time of maximal IkBα degradation after a ligand challenge given at tₓ (Fig. 1C); for a schematic timeline, see Fig. 2B). IkBα-FLuc reporter levels in these lysates were analyzed by bioluminescence imaging (upon the addition of saturating D-luciferin and ATP), and endogenous IkBα levels were determined by Western blot analysis and semiquantitative densitometric analysis (Fig. 2C). From these data, we were then able to calculate responsiveness levels for both IkBα and IkBα-FLuc as a function of time after TNFα preconditioning. Responsiveness at each challenge time was calculated by determining the magnitude of IkBα degradation induced by TNFα challenge divided by the magnitude of IkBα degradation in unpreconditioned cells from the same plate. Specifically, the ratio at tₓ + 25 min of IkBα in preconditioned cells challenged with TNFα over preconditioned cells challenged with vehicle was divided by the ratio at tₓ + 25 min of IkBα in unpreconditioned cells challenged...
with TNFα over unpreconditioned cells challenged with vehicle, the latter ratio representing the maximal possible response. We observed a strong correlation ($r = 0.95$) between levels of responsiveness for endogenous IκBα and IκBα-FLuc (Table 1). Consistent with our earlier observations derived from live cell dynamic bioluminescence imaging experiments (Fig. 2A), we observed that at 30 min postpreconditioning, cells were approximately half as responsive as naive (i.e. unpreconditioned) cells to a TNFα challenge and had gained full responsiveness by 120 min. Thus, a transient refractory period seemed to exist from 30 to 120 min post-TNFα treatment. Data are presented as -fold initial, -fold TNFα—Hypothetically, this loss and regain of the capacity of cells to process IκBα can be explained by 1) internalization or shedding of TNFα receptors, followed by their recycling to the cell membrane (26, 27), 2) transient down-regulation of IKK activity as previously reported (15, 28), or, alternatively, 3) by a yet unknown mechanism of regulation, downstream of IKK. We therefore sought to establish the relative contributions of receptor dynamics and IKK regulation to this refractory period.

To determine the extent of receptor dynamics in governing the observed loss and regain of IκBα processing, we took advantage of a discovery, made 20 years ago (14), that IL-1β induces transient down-regulation of TNFα receptors but not vice versa (i.e. TNFα has no effect on either the affinity or the number of IL-1β surface receptors), as tested in a variety of cell lines and primary cells. Hence, we aimed to determine IκBα responsiveness to an IL-1β challenge as a function of time after TNFα preconditioning in HepG2 cells. Cells expressing pxB5→IκBα-FLuc were treated with a 30-s pulse of TNFα (20 ng/ml), followed by a continuous challenge with IL-1β, initiated at increasing P-C intervals (0–240 min). IκBα processing was analyzed

FIGURE 2. TNFα-induced a transient refractory period for IκBα processing. A, dynamic live cell bioluminescence imaging profiles of IκBα-FLuc from TNFα preconditioning plus challenge experiments. The black arrows denote 30-s preconditioning pulse; red arrows denote the beginning of continuous TNFα challenge; black profiles represent cells preconditioned and then challenged at the indicated time points; red profiles represent cells treated at time 0 with continuous TNFα (denoting the maximal possible degradation response of IκBα upon continuous TNFα treatment). Data are presented as -fold initial, -fold TNFα-untreated. B, schematic representation of the experimental timeline as used in C. Cells were preconditioned with TNFα for 30 s and then, at increasing intervals (0–240 min), were continuously challenged with TNFα. The arrowheads represent when cells were harvested and lysates were prepared (25 min post-challenge for quantitative bioluminescence imaging and Western blot analysis). C, IκBα-FLuc and endogenous IκBα levels, 25 min post-TNFα or vehicle challenge, as measured by bioluminescence imaging and Western blot, respectively.

TABLE 1

Percentage responsiveness of IκBα processing

Quantification of IκBα-FLuc and IκBα responsiveness to a second continuous challenge of TNFα at the indicated interval following a 30-s preconditioning pulse of TNFα was determined from the bioluminescence imaging and Western blot data shown in Fig. 2C. Responsiveness at each challenge time was calculated by determining the percentage of challenge-specific IκBα degradation divided by the percentage of IκBα degradation in unpreconditioned cells from the same plate. The responsiveness of IKK was determined by an IKK kinase assay.

| Responsiveness of IκBα processing | 30 min after TNFα preconditioning | 60 min after TNFα preconditioning | 120 min after TNFα preconditioning | 240 min after TNFα preconditioning |
|-----------------------------------|----------------------------------|-----------------------------------|-----------------------------------|-----------------------------------|
| IκBα (Western blot)              | 49                               | 80                                | 100                               | 82                                |
| IκBα-FLuc (bioluminescence)      | 43                               | 74                                | 90                                | 94                                |
| IKK activity                     | 35                               | 56                                | 75                                | 69                                |
Refractory Period in NF-κB Signaling

by live cell dynamic bioluminescence imaging (Fig. 3A). Using this experimental setup, we again observed a transient refractory period (from 30 to 120 min post-TNFα preconditioning) during which HepG2 cells exhibited decreased IkBα responsiveness. The magnitude of the ligand-induced degradation increased as the interval to the IL-1β challenge increased, becoming fully responsive again by 120 min (Fig. 3A). These data suggested that even in the absence of ligand-induced receptor desensitization or cross-regulation, the capacity of cells to process IkBα was compromised within the first 2 h after a short TNFα stimulation.

We next aimed at deciphering whether transient down-regulation of IKK activity could explain the observed loss and regain in IkBα responsiveness. We therefore performed a series of IKK kinase assays in order to directly measure the temporal activity profile of IKK, a central junction of the TNFα and IL-1β pathways that integrates signals from a myriad of upstream regulators (e.g. TNF receptor-associated factors, mitogen-activated protein/extracellular signaling-regulated kinase kinase, TGFβ-activated kinase-binding protein, TGFβ-activated kinase, NF-κB-inducing kinase, receptor-interacting protein, A20, protein kinase Cζ, etc. (2, 7, 29)). HepG2 cells were treated with TNFα (20 ng/ml) either as a 30-s pulse or continuously. At the indicated time points, cells were harvested, and IKK complexes were immunoprecipitated and assayed for their capacity to phosphorylate exogenous GST-IκBα-(1–54) (23). We found that for both 30-s pulses and continuous TNFα exposure, temporal profiles of IKK activity were almost identical, with both peaking at 10 min. However, consistent with our earlier findings that continuous TNFα treatment elicits greater IkBα degradation than a 30-s pulse (Fig. 1C), continuous TNFα treatment exhibited slightly elevated and more sustained levels of IKK activity compared with pulsed TNFα treatment (Fig. 3B). Importantly, Western blot analysis showed that IKK complex levels (as determined by IKKα protein) did not change over the experimental time course (Fig. 3C), confirming that the increase in net kinase activity was due specifically to IKK activation.

IKK-KA data were also collected from preconditioned cells, 10 min post-challenge (at the time of maximal IKK activity; see Fig. 3B) at increasing P-C intervals (0–240 min). Using these data together with the IKK activity profiles generated for 30-s pulse and continuous TNFα treatment regimens (Fig. 3B), we were able to calculate the net capacity of IKK to phosphorylate IkBα as a function of time after TNFα preconditioning (i.e. IKK responsiveness (Table 1); see “Experimental Procedures” for details on this calculation). Based on this calculation, we noted that the capacity of IKK to respond to a second challenge of TNFα was significantly compromised at 30 min post-TNFα preconditioning and then gradually increased, reaching ~75% responsiveness by 120 min. Up to 240 min, IKK activity did not fully recover to initial levels, consistent with other reports indicating that upon TNFα stimulation, IKK activity rapidly and transiently declines due to autocatalytic C-terminal hyperphosphorylation (15) and protein phosphatase 2C-dependent dephosphorylation (16), followed by late NF-κB-dependent down-regulation, a process attributed, in part, to A20, an IKK-inhibitory protein (29). Hence, these data suggested that 1) the observed ligand-induced transient refractory period of IkBα processing (Figs. 2 and 3 and Table 1) correlated only in part with ligand-induced transient down-regulation of IKK activity and that 2) the level to which cells are able to degrade IkBα was not linear with the capacity of IKK to phosphorylate IkBα (i.e. full IkBα responsiveness was observed as soon as 120 min post-TNFα preconditioning (Figs. 2 and 3A), a time point where IKK responsiveness was still compromised (Table 1)). These data indicated that either sub-maximal IKK activity could now fully support ligand-induced IkBα degradation following the refractory period or that additional ligand-responsive elements existed that converged on IkBα to induce a full response.
Refractory Period in NF-κB Signaling

FIGURE 4. Computational simulation of IκBα responsiveness in the absence of upstream receptor or IKK regulation. A, interpolated and extrapolated (0–360 min, at 5-min intervals) IKK activity profiles (right) of cells treated continuously (C, green curve) or by a 30-s pulse (P, blue curve) of TNFα (20 ng/ml) were used as inputs to computationally simulate total IκBα dynamics (right). B–E, left panels, hypothetical IKK activity profiles of preconditioned cells, challenged at the indicated times (denoted by black arrowheads) with a second, continuous dose of TNFα were generated by superimposing the continuous TNFα-induced IKK activity profiles at increasing intervals after the 30-s pulse TNFα-induced IKK profile. For generating these hypothetical profiles, we assume no precondition-induced receptor or IKK regulation. Right panels, the hypothetical IKK profiles were used as inputs into the model to predict IκBα dynamics. Note that challenge-induced IκBα degradation (initiated at the red arrowhead) is recovered in a time-dependent manner.

Computational Modeling of NF-κB Signaling Suggested an Additional Layer of Regulation, Downstream of IKK, Governing the Observed Refractory Period for IκBα Processing—The NF-κB pathway provides an excellent example of a complex signaling system employing numerous temporally distinct autoregulatory mechanisms and negative feedback loops. IKK enzymatic activity, which is both endogenously and exogenously regulated, controls the degradation of its own substrate (IκBα), which is later strongly up-regulated in an NF-κB-dependent manner (Fig. 1A). Rapid changes in substrate availability, conformation, and subcellular localization imply that alternative mechanisms of regulation might exist other than changes in enzymatic activity. Although a ligand-induced transient refractory period of IκBα processing could be explained in part by down-regulation of IKK activity, we were intrigued to examine whether an alternative regulatory mechanism, based on substrate (IκBα) dynamics, might exist to complement or “back up” IKK regulation. Obviously, inhibition of IKK was not a viable option for analyzing downstream regulation, since such inhibition will result in complete loss of responsiveness in the absence or presence of preconditioning. We therefore decided to undertake a computational approach and explore IκBα dynamics in silico, assuming no down-regulation of IKK activity. We used a well accepted computational model that used experimentally or hypothetically driven IKK activity profiles as inputs and, in return, calculated ligand-induced dynamics of 24 different subpopulations of mediators on the IKK-NF-κB axis.

As a first step, to test the robustness of the model, we sought to compare our IκBα-FLuc bioluminescence imaging data for 30-s pulsing and continuous TNFα treatments (Fig. 1C) with the dynamics of IκBα, as predicted by the computational model. To accomplish this, we used as inputs the IKK activity profiles generated for 30-s pulse and continuous TNFα treatment regimens (Fig. 4A, left; see “Experimental Procedures” for details on numerical processing of the raw data to fit the model). The dynamics of six different free and complexed IκBα subpopulations could be predicted by the model (i.e. free IκBαcyt, IκBαIKKcyt, IκBαNF-κBcyt, IκBαIKK-NF-κBcyt, free IκBαnuc, and IκBαNF-κBnuc). Since live cell bioluminescence imaging of IκBα-FLuc could not distinguish between these populations, we summed up the predicted concentrations of all IκBα subpopulations and plotted the predicted total IκBα levels as a function of time (Fig. 4A, right). For both treatment regimens, we noted an excellent correlation between the predicted profiles of IκBα and the experimentally generated profiles of IκBα-FLuc (Fig. 1C). The timing and extent of IκBα degradation as well as the overall dynamic behavior were highly similar. However, differences in the amplitude and timing of resynthesis (experimental: ~8-fold initial at ~120 min; computational: 1.2–1.5-fold initial at ~90 min) were observed and could be explained by dynamic differences between the endogenous IκBα promoter and the synthetic κB5-TATA promoter driving IκBα-FLuc (i.e. differences in binding affinity and cooperativity toward NF-κB).

We next generated hypothetical IKK profiles representing IKK activities from preconditioned/challenged cells, assuming no upstream receptor or IKK regulation (i.e. experimentally derived challenge-specific IKK activity was overlaid on top of experimentally derived precondition-specific residual IKK activity). These hypothetical IKK activity profiles (Fig. 4, B–E, left, each generated with a different P-C interval) were used as inputs for computing total IκBα dynamics (Fig. 4, B–E, right). Surprisingly, the computational model predicted that even in the absence of receptor dynamics or IKK regulation, IκBα processing would be transiently compromised (compare, for example, the second, challenge-induced degradation phase at 120 or 240 min with the ones at 30 or 60 min). These data suggested that although IKK down-regulation partially correlated with the ligand-induced transient refractory period for IκBα processing, an additional regulatory mechanism was present downstream of IKK. Importantly, IκBα availability per se was not
sufficient to explain changes in IκBα responsiveness, because, as confirmed experimentally and computationally, at 60 min post-preconditioning, the IκBα concentration had already recovered, whereas degradation potential was still low (compare Fig. 2, A and C, Table 1, and Fig. 4C).

**Nuclear Export of IκBα—NF-κB complexes may also control the capacity of cells to process IκBα.** Having demonstrated experimentally the phenomenon of a ligand-induced transient refractory period for IκBα processing and after dissecting biochemically and computationally the origins of this observation, we next sought to more closely examine the components of the computational model in order to identify candidates, down-stream of IKK, capable of regulating IκBα responsiveness. While examining the rate constants of a variety of reactions used by the model, we noticed that free versus NF-κB-bound IκBα differed tremendously in their capacity to associate with IKK (1.35 versus 11.1 μM⁻¹ min⁻¹, respectively) and to be degraded in an IKK-dependant manner (0.12 versus 0.00006 min⁻¹, respectively). These differences in IKK association and ligand-induced degradation were experimentally established by Zandi et al. (30).

This led us to put forward the following model (Fig. 5A). 1) free IκBα and NF-κB-bound IκBα represent “protected” and “unprotected” populations with respect to ligand-induced, IKK-dependent proteasomal degradation. 2) Under steady-state conditions, there is a stoichiometric excess of IκBα over NF-κB in the cytoplasm (~0.7 NF-κB per IκBα according to the model). This may explain our observations that even at saturating concentrations of TNFα or IL-1β, IκBα degradation never exceeded 70–80% of initial level (e.g. Fig. 1C). 3) Upon ligand stimulation, NF-κB-bound IκBα is degraded, NF-κB translocates to the nucleus, and IκBα is resynthesized. 4) At this point, although IκBα is highly abundant, its capacity to be degraded in response to a second stimulus is still severely compromised, because NF-κB is in the nucleus. 5) IκBα can freely shuttle between the cytoplasm and the nucleus, pulling NF-κB molecules (that lack nuclear export signals (31)) back to the cytoplasm. This step may be the rate-limiting step for acquisition of full responsiveness. 6) Newly synthesized IκBα molecules uncomplexed with NF-κB are rapidly degraded (32), and only after all NF-κB molecules are recovered back to the cytosol and the NF-κB-bound IκBα over free IκBα ratio returns to pre-stimulation levels (~0.7) are cells able to mount a full response again.

To experimentally examine the nuclear export hypothesis, we sought to analyze ligand-induced changes in cytoplasmic IκBα-NF-κB complexes. However, the computational model predicted that ligand-induced changes of cytoplasmic IκBα-NF-κB and total cytoplasmic NF-κB were essentially the same (i.e. at any given time, virtually all cytoplasmic NF-κB was bound to IκBα; Fig. 5B), suggesting that monitoring cytoplasmic total NF-κB was an excellent approximation for following cytoplasmic IκBα-NF-κB complexes. We therefore pulsed HepG2 cells for 30 s with TNFα (20 ng/ml), and at various times after stimulation, we fixed, permeabilized, and immunostained the cells for p65 NF-κB (Fig. 5C). We found that upon a 30-s TNFα pulse, p65 rapidly translocated to the nucleus (maximal by 30 min) but by 60–120 min was back in the cytoplasm. The excellent temporal correlation between the levels of cytoplasmic NF-κB (as derived computationally or experimentally; Fig. 5, B and C, respectively) and the competence of cells to degrade IκBα in response to a proinflammatory ligand (i.e. Table 1) strongly suggested that nuclear transport of NF-κB provided a potential alternative mechanism to transiently desensitize IκBα processing (refractory period), in addition to the mechanism of IKK down-regulation (Fig. 3, B and C, and Table 1).

**DISCUSSION**

Ligand-induced desensitization is a common theme in many biological systems (13), thereby allowing cells to mount an appropriate response independently of ligand exposure time. Thus, prolonged exposures will not result in excessive responses, but instead, cells are enabled to build up a downstream response while being unable to perceive a second activating cue. Desensitization and resensitization are traditionally perceived to be linked to receptor dynamics (internalization, shedding, and recycling); however, any mediator or regulator along a signaling pathway can be hypothetically desensitized, therefore transiently blocking signal transduction (13).

In this work, we demonstrated that although cells can efficiently activate NF-κB in response to a TNFα exposure as short as 30 s, such stimulation was followed by a refractory period during which the capacity of cells to respond to a second homologous or heterologous stimulus was severely compromised. We further found that this transient refractory period correlated in part with a temporal down-regulation of IKK activity but not with receptor desensitization. Computational modeling enabled us to identify an additional layer of regulation, downstream of IKK, controlling the capacity of cells to respond to a second challenge. Ligand-induced dynamic changes in substrate (IκBα) availability, conformation, and subcellular localization form the basis for this mechanism. Further analysis led us to conclude that nuclear export of NF-κB may be a rate-limiting step in controlling IκBα homeostatic metabolism, a term recently coined by O’Dea et al. (33).

Our study highlights the multifaceted regulation of NF-κB signaling (Fig. 6) and sheds light on the refractory nature of IκBα processing as a route to transiently desensitize NF-κB activity upon subsequent rounds of stimulation. Rapid and transient deactivation of IKK activity as well as temporal reduction in its capacity to respond to a subsequent challenge (IKK responsiveness) seems to play a crucial role in this process. Previous studies indicated that both the amplitude and the timing of IKK activation affect not only the intensity of NF-κB-dependent transcription but also the specificity of the transcriptional response (23, 34). This indicated that besides resolution of the inflammatory response and induction of a refractory period (temporally preventing subsequent rounds of IκBα degradation upon restimulation), rapid down-regulation of IKK activity (28) plays a pivotal role in determining the type of elicited transcriptional program.

In addition to IKK regulation, our work demonstrated that nuclear export of IκBα-NF-κB complexes may have also regu-
Refractory Period in NF-κB Signaling

In the present and previous studies (20), we demonstrated that dynamic bioluminescence imaging of IκBα-Fluc reporters in live cells provides robust and accurate readouts of ligand-induced IκBα dynamics. In effect, real time bioluminescence imaging was equivalent to performing continuous on-line Western blots of IκBα at 5-min intervals. An analogous transcriptionally coupled reporter (κB₅−>IκBα-EGFP) was generated by Nelson et al. (35) for monitoring IκBα dynamics in single cells by live cell fluorescence microscopy. Although such a system provides the means to monitor ligand-induced translocations and oscillations in IκBα levels, temporal resolution of this reporter is limited by the long maturation time of EGFP (>1 h) (38, 39). This notion and the fact that Nelson et al. (35, 36) co-overexpressed p65-red fluorescent protein may explain the vast difference between the observed period of IκBα-EGFP oscillations (~300 min) and the period of endogenous IκBα oscillations, as predicted computationally (~90–120 min) (25).

Although longer term IκBα oscillatory behavior was not the focus of the present study, we did observe single oscillations within 150–180 min. Because FLuc is active immediately upon translation, our reporter should afford greater temporal resolution, enabling accurate readouts of IκBα dynamics and oscillations in live cells for such studies as well as the multistimulation protocols as described herein.

Of note, a previous study aimed at analysis of IκBα stabilization indicated a role for p38 in IκBα stabilization and, in some cell lines, in prevention of sequential degradation of IκBα upon concurrent exposure to TNFα following continuous pretreatment with IL-1β (40). However, since IL-1β has been shown to induce rapid and dramatic down-regulation of TNFα receptors (but not vice versa) (14), inhibition of TNFα-induced IκBα processing, as observed by Place et al. (40), could be attributed directly to receptor dynamics rather than IκBα stabilization. This confounding factor highlights the importance of asymmetric receptor cross-desensitization, a phenomenon that remains poorly understood but has far reaching physiological consequences.

In conclusion, TNFα preconditioning protocols and dynamic imaging revealed a transient suppression of the capacity of cells to process IκBα. This refractory period for IκBα processing was controlled both by IKK activity and NF-κB distribution. In particular, the data suggested that nuclear export of NF-κB may provide additional rate-limiting regulation governing the refractory period machinery. These regulatory mechanisms provide a “molecular timer” controlling the amplification in turnover rate. Following the peak of IκBα resynthesis, both endogenous IκBα and our IκBα-Fluc reporter begin returning to base-line levels faster under continuous TNFα treatment, suggesting that ligand-induced reactivation of IκBα degradation is occurring under continuous TNFα exposure, as expected (23).

In the present study, we also observed that the endogenous promoter contains three distant NF-κB sites, whereas the synthetic promoter contains five tandem high affinity κB sites. Nevertheless, since both endogenous IκBα and IκBα-Fluc exhibit similar half-life times (20), differences in the timing of resynthesis cannot be explained by differences in turnover rate. Following the peak of IκBα resynthesis, both endogenous IκBα and our IκBα-Fluc reporter begin returning to base-line levels faster under continuous TNFα treatment, suggesting that ligand-induced reactivation of IκBα degradation is occurring under continuous TNFα exposure, as expected (23).

In the present study, we also observed that the endogenous promoter contains three distant NF-κB sites, whereas the synthetic promoter contains five tandem high affinity κB sites. Nevertheless, since both endogenous IκBα and IκBα-Fluc exhibit similar half-life times (20), differences in the timing of resynthesis cannot be explained by differences in turnover rate. Following the peak of IκBα resynthesis, both endogenous IκBα and our IκBα-Fluc reporter begin returning to base-line levels faster under continuous TNFα treatment, suggesting that ligand-induced reactivation of IκBα degradation is occurring under continuous TNFα exposure, as expected (23).
Refractory Period in NF-κB Signaling

titude, timing, and specificity of the NF-κB-mediated transcriptional program.

Acknowledgments—We thank Prof. Alexander Hoffmann, Shannon Werner, and Derren Barken (University of California, San Diego) for providing the computational model software. We also thank Dr. Yun-Feng Feng and Prof. Greg Longmore (Washington University) and Prof. Phillip Cohen (University of Dundee) for advice with the IKK kinase assays and Dr. Dustin Maxwell (Washington University) for assistance with immunofluorescence microscopy.

REFERENCES
1. Han, J., and Ulevitch, R. J. (2005) Nat. Immunol. 6, 1198–1205
2. Winsauer, G., and de Martin, R. (2007) Thromb. Haemostasis 97, 364–369
3. Hoffmann, A., and Baltimore, D. (2006) ImmunoL Rev. 210, 171–186
4. Perkins, N. D. (2007) Nat. Rev. Mol. Cell. Biol. 8, 49–62
5. Karin, M., and Greten, F. R. (2005) Nat. Rev. Immunol. 5, 441–446
6. Karin, M. (2006) Science 312, 315–318
7. Keyse, S. M., and Ghosh, S. (2004) Nat. Rev. Mol. Cell. Biol. 5, 53–67
8. Bunting, K., Rao, S., Hardy, K., Woltring, D., Denyer, G. S., Wang, J., Naamane, N., van Helden, J., and Eizirik, D. L. (2007) BMC Bioinformatics 8, 55
9. Kim, H. J., Hawke, N., and Baldwin, A. S. (2006) Cell Death Differ. 13, 738–747
10. Karin, M., Cao, Y., Greten, F. R., and Li, Z. W. (2002) Nat. Rev. Cancer 2, 301–310
11. Karmann, K., Min, W., Fanslow, W. C., and Pober, J. S. (1996) J. Exp. Med. 184, 173–182
12. Alberts, B., Bray, D., Lewis, J., Raff, M., Roberts, K., and Watson, J. D. (1994) in Molecular Biology of the Cell, 3rd Ed., pp. 771–785, Garland Publishing, New York
13. Delhase, M., Hayakawa, M., Chen, Y., and Karin, M. (1999) Science 284, 309–313
14. Prajapati, S., Verma, U., Yamamoto, Y., Kwak, Y. T., and Gaynor, R. B. (2004) J. Biol. Chem. 279, 1739–1746
15. Trinchieri, G., and Sherry, S. (2007) Nat. Rev. Immunol. 7, 179–190
16. Gloire, G., Legrand-Poels, S., and Piette, J. (2006) Biochem. Pharmacol. 72, 1493–1505
17. Blanque, R., Meakin, C., Millet, S., and Gardner, C. R. (1998) Gen. Pharmacol. 31, 301–306
18. Gross, S., and Pignoni-Worms, D. (2005) Nat. Methods 2, 607–614
19. Nejari, M., Hafdi, Z., Dumortier, J., Bringui, A. F., Feldmann, G., and Scoazec, J. Y. (1999) Int. J. Cancer 83, 518–525
20. Wullaert, A., van Loo, G., Heyninck, K., and Beyaert, R. (2007) Endocr. Rev. 28, 365–386
21. Werner, S. L., Barken, D., and Hoffmann, A. (2005) Science 309, 1857–1861
22. Hastie, C. J., McLauchlan, H. J., and Cohen, P. (2006) Nat. Protoc. 1, 968–971
23. Hoffmann, A., Levchenko, A., Scott, M. L., and Baltimore, D. (2002) Science 298, 1241–1245
24. Higuchi, M., and Aggarwal, B. B. (1994) J. Immunol. 152, 3550–3558
25. Dri, P., Gasparini, C., Menegazzi, R., Cramer, R., Alberi, L., Presani, G., Garbisa, S., and Patriarca, P. (2000) J. Immunol. 165, 2165–2172
26. Cheong, R., Krelle, M., Werner, S. L., Regal, J., Hoffmann, A., and Levchenko, A. (2006) J. Biol. Chem. 281, 2945–2950
27. Liu, Y. C., Penninger, J., and Karin, M. (2005) Nat. Rev. Immunol. 5, 941–952
28. Zandi, E., Chen, Y., and Karin, M. (1998) Science 281, 1360–1363
29. Huxford, T., Huang, D. B., Malek, S., and Ghosh, G. (1998) Cell 95, 759–770
30. Pando, M. P., and Verma, I. M. (2000) J. Biol. Chem. 275, 21278–21286
31. O’Dea, E. L., Barken, D., Peralta, R. Q., Tran, K. T., Werner, S. L., Kears, J. D., Levchenko, A., and Hoffmann, A. (2007) Mol. Syst. Biol. 3, 111
32. Covert, M. W., Leung, T. H., Gaston, J. E., and Baltimore, D. (2005) Science 309, 1854–1857
33. Nelson, D. E., Ikekwaba, A. E., Elliott, M., Johnson, J. R., Gibney, C. A., Foreman, B. E., Nelson, G., See, V., Horton, C. A., Spiller, D. G., Edwards, S. W., McDowell, H. P., Unitt, J. F., Sullivan, E., Grimley, R., Benson, N., Broomhead, D., Kell, D. B., and White, M. R. (2004) Science 306, 704–708
34. Barken, D., Wang, C. J., Kears, J., Cheong, R., Hoffmann, A., and Levchenko, A. (2005) Science 308, 52
35. Kears, N. S., and Derren Barken (2006) J. Cell Biol. 173, 659–664
36. Sniegowski, J. A., Lappe, H. N., Patel, H. N., Huffman, H. A., and Wachter, R. M. (2005) J. Biol. Chem. 280, 26248–26255
37. Zhang, L., Patel, H. N., Lappe, J. W., and Wachter, R. M. (2006) J. Am. Chem. Soc. 128, 4766–4772
38. Place, R. F., Haspeslagh, D., and Giardina, C. (2003) J. Cell. Physiol. 195, 470–478