Signaling via the NFkB system



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The nuclear factor kappa B (NF κ B) family of transcription factors is a key regulator of immune development, immune responses, inflammation, and cancer. The NFkB signaling system (defined by the interactions between NFkB dimers, IkB regulators, and IKK complexes) is responsive to a number of stimuli, and upon ligand-receptor engagement, distinct cellular outcomes, appropriate to the specific signal received, are set into motion. After almost three decades of study, many signaling mechanisms are well understood, rendering them amenable to mathematical modeling, which can reveal deeper insights about the regulatory design principles. While other reviews have focused on upstream, receptor proximal signaling (Hayden MS, Ghosh S. Signaling to NF-KB. Genes Dev 2004, 18:2195-2224; Verstrepen L, Bekaert T, Chau TL, Tavernier J, Chariot A, Bevaert R. TLR-4, IL-1R and TNF-R signaling to NF-κB: variations on a common theme. Cell Mol Life Sci 2008, 65:2964–2978), and advances through computational modeling (Basak S, Behar M, Hoffmann A. Lessons from mathematically modeling the NF-kB pathway. Immunol Rev 2012, 246:221–238; Williams R, Timmis J, Owarnstrom E. Computational models of the NF-KB signalling pathway. Computation 2014, 2:131), in this review we aim to summarize the current understanding of the NFκB signaling system itself, the molecular mechanisms, and systems properties that are key to its diverse biological functions, and we discuss remaining questions in the field. © 2016 Wiley Periodicals, Inc.

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INTRODUCTION

Nuclear factor kappa B (NF κ B) is a family of dimeric transcription factors central to coordinating inflammatory responses; innate and adaptive immunity; and cellular differentiation, proliferation, and survival in almost all multicellular organisms.^{1–4} The NF κ B system is tightly regulated, and misregulation of NF κ B has been implicated in a wide range of diseases ranging from cancers to inflammatory and immune disorders. As a result, the NF κ B regulatory network and its dynamics offer a multitude of promising therapeutic targets that remain to be fully explored and translated into clinical use.^{5–7} However, there continues to be an untapped potential for finer grained therapeutic targeting of the NF κ B signaling system that requires a quantitative understanding of dynamical control and the integration of various physiological and pathological signals and stimuli.^{8–10}

In Mammalia, the NFkB network consists of five family member protein monomers (p65/RelA, RelB, cRel, p50, and p52) that form homodimers or heterodimers that bind DNA differentially¹¹⁻¹⁴ and are regulated by two pathways: the canonical, NFKB essential modulator (NEMO)-dependent pathway and the noncanonical, NEMO-independent pathway. These pathways tightly control the levels and dynamics of the transcriptionally active NFkB dimer repertoire constitutively and in response to stimuli, and thus control broad gene expression programs^{15,16} via the recruitment of co-activators¹⁷ or interplay with other transcription factors.^{18,19} The activation pathways control NFkB activity through multiple mechanisms: degradation of IkB inhibitor proteins, processing of NFkB precursor proteins, and expression of NFkB monomer proteins.^{10,20} Signals from

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tumor necrosis factor receptor (TNFR), toll-like receptor (TLR) superfamilies, interleukin receptor (IL-1R) and metabolic genotoxic, and shear stresses are integrated by the IkB/NFkB signaling network to produce signal-specific, context-specific, and celltype-specific transcriptional responses.^{21–23}

CANONICAL SIGNALING

Signaling via the NEMO-Associated IKK Complex

The canonical NFkB signaling pathway (a.k.a. NEMO-dependent pathway) is mediated by kinase complexes consisting of the scaffold/adaptor protein NEMO (a.k.a. IKK γ) and two I κ B kinases (IKK1/2, a.k.a. IKK α and IKK β) (Figure 1(a)). This IKK complex is activated by mechanisms that are NEMO dependent. The IKK kinases are activated by phosphorylation of serines in the activation T-loop, characteristic of the MAPK superfamily, and three mechanisms for IKK activation have emerged: (1) NEMO multimerizes IKK subunits²⁷ to allow for activation via trans-autophosphorylation, (2) and/or brings them in proximity to upstream kinases such as TAK1,²⁸ which may also lead to mutual activation forming positive feedback and digital dose-response characteristics.^{29,30} These mechanisms are mediated by NEMO's ubiquitin-binding domain that allows for IKK's recruitment to nondegradative K63-linked ubiquitin chains, which are a hallmark of inflammatory signaling. Finally, (3) NEMO itself is a substrate of ubiquitination, particularly linear ubiquitin chains produced by the LUBAC enzyme,³¹ which also facilitates the formation of transient signalsomes.

A wide variety of inflammatory cytokines (such as TNF and IL-1), a wide variety of pathogenassociated molecular patterns (PAMPs), or antigen/ immune stimulatory signals result in IKK phosphorylation-dependent activation of the NEMOcontaining complex, by one or a subset of these mechanisms,²⁴ resulting in complex dynamical control.³² Once activated, the complex binds to and phosphorylates IkB proteins on specific serines in the N-terminal, leading to ubiquitination and subsequent proteasomal degradation.²⁵ The degradation of inhibitors releases NFkB dimers associated with them, freeing NFkB dimers and allowing them to bind kB site-containing DNA and thus rapidly accumulate in the nucleus. Following IkB release the NFkB subunits are subject to a variety of posttranslational modification that fine-tune gene-expression control.³³

Further, NEMO was shown to function as a scaffold between IKK and $I\kappa B\alpha$, thereby directing



FIGURE 1 | The canonical nuclear factor κ B (NF κ B) activation pathway. (a) Schematic depiction of the canonical NF κ B signaling pathway. Multiple inflammatory signals activate the complex containing NEMO and IKK1/2. IKK1/2 phosphorylates NF κ B-bound I κ Bs, targeting them for ubiquitination and proteasomal degradation.^{24,25} Free I κ Bs also undergo constitutive degradation via a ubiquitin-independent proteasomal degradation pathway. As I κ Bs are degraded, free NF κ B is then able to translocate to the nucleus where it binds to κ B sites on DNA and activates gene expression.^{21,22} I κ B α , β , and ε are themselves NF κ B target genes, along with p100 that can form higher-molecular-weight complexes that inhibit NF κ B.^{25,26} (b) Diagram of the regulatory logic of the canonical NF κ B signaling network. Canonical signals activate NEMO/IKK, downregulating I κ Bs and reducing inhibition of NF κ B. Free NF κ B then translocates to the nucleus where it upregulates I κ Bs and p100 and in turn I κ Bô.

IKK activity to $I\kappa B\alpha$.³⁴ This mechanism ensures that the activation of NF κ B dimers associated with $I\kappa B\alpha$, which is RelA:p50 in most cells and conditions, is directly linked to signals propagating through the NEMO hub. Hence, canonical signaling is often thought to be synonymous with RelA activation, but this is not always the case. In inflammatory dendritic cells, $I\kappa B\alpha$ was also shown to be associated with RelB:p50 dimers, thus rendering RelB a key transcriptional effector of the canonical NFκB pathway

in that cell type.³⁵ While the activation mechanism of IKK is beginning to be elucidated with the aid of recent structural and biophysical characterizations^{36,37} and, for example, combined single-cell computational studies that have identified distinct pathways for robust digital responses and noisy sustained responses,³⁸ how IKK is inactivated remains unclear. One attractive proposal is that IKK is regulated in an autocatalytic cycle of at least three states in which activation occurs from a poised state and is followed by an inactive state. While the dynamic properties of such a kinase control cycle have been studied,³⁹ the biophysical evidence remains scant, but could involve trans-autophosphorylation of an inhibitory C-terminal domain in IKK640 and/or conformational changes of the complex.⁴¹

IκBα Negative Feedback

Among the target genes regulated by κB sites are the IkBs that, upon transcriptional induction and resynthesis, are able to translocate to the nucleus, bind to and inhibit NFkB activity, trafficking it back to the cytosol. This constitutes the primary component of the self-regulating negative feedback loop²⁵ (Figure 1(b)). This feedback loop not only prevents run-away NFkB activity in response to transient inflammatory signals but also poises the system for reactivation when IKK activity is longer lasting. IkBa negative feedback has been studied in some detail with a combined experimental and mathematical modeling approach, and interesting properties have emerged: (1) Given the delay intrinsic to $I\kappa B\alpha$ resynthesis, even very transient cytokine exposure (1 min TNF) results in almost a full hour of NFkB activity.⁴² That 1 h of NFkB activity is invariant to the duration of the signal unless the incoming signal lasts longer than approximately 45 min. (2) Transcriptional induction of IkBa is necessary but not sufficient for mediating this effective negative feedback control of NF κ B activity.⁴³ I κ B β , when controlled by an NFkB-inducible promoter, is unable to provide normal physiologically observed negative feedback⁴⁴; nuclear import, export, and protein degradation mechanisms specific to $I\kappa B\alpha$ are critically important to recapitulating proper negative feedback and the NFkB dynamic responses characteristic of normal activity.^{45–49} In addition, $I\kappa B\alpha$ has been shown to be able to strip NFkB off DNA (or associate with chromatin⁵⁰), a function that $I\kappa B\beta$ does not possess.⁵¹ (3) Given effective $I\kappa B\alpha$ negative feedback,

stimulation conditions that produce longer lasting IKK activities allow for repeated cycles of reactivation, leading to oscillatory NF κ B activity observed in biochemical bulk population assays²⁵ or single-cell assays.^{38,52–54} Intrinsic variability in the kinetic mechanisms that govern I κ B α feedback is thought to render the oscillatory behavior more robust to variations in dynamic signals.^{55,56} However, the potential function of NF κ B oscillations remains unclear, and no satisfying answer has been presented as to how such oscillations are interpreted by downstream gene regulatory networks.⁵⁷

IκBδ Negative Feedback

Another transcriptional target gene that is induced by nuclear NFkB activity is the nfkb2 gene, which produces the p100 protein.⁴ p100 was first known as the precursor for the p52 protein, a dimerization partner of RelB and potentially other Rel proteins. However, p100 that is not processed to p52 is able to form higher-molecular-weight complexes that are capable of binding to NFkB, acting as an IkB, termed IkBô.²⁶ As such, NFkB control of nfkb2 forms another negative feedback loop that may terminate NFkB signaling.58 However, transcriptional induction and protein synthesis are slow, in part due to the length and half-life of the mRNA and protein, and the subsequent required oligomerization step also takes several hours.⁵⁹ Thus, in contrast to $I\kappa B\alpha$ that functions rapidly, IkBô's role is primarily in attenuating persistent signals.58 Indeed, while IkBa negative feedback is reversible as it is a sensitive substrate for continued canonical IKK activity, IkBo is insensitive to canonical signals and thus attenuates the canonical pathway regardless of whether incoming signals persist. Further, because IkB8 has a longer half-life than other IkBs, it contributes to a signaling memory in which sequential stimulation events are dampened.⁵⁸ However, as a substrate for noncanonical signaling (see below), IkBo is a signaling crosstalk node that integrates canonical and noncanonical signals that may result in noncanonical signals emanating from $LT\beta R$ or BAFF to activate (or prolong the activation of) NFκB RelA or cRel dimers.^{26,60,61}

Other Feedback Mechanisms

There are several other negative and positive feedback mechanisms that contribute to the complex and potentially oscillatory dynamics of nuclear NF κ B.^{58,62} I κ Be was shown to provide delayed negative feedback (due to a transcriptional delay) that may partially compensate for the loss of I κ B α ,⁶³ and was suggested to dampen IkB α -mediated oscillations by forming a dual, antiphase negative feedback system.⁶² There is also evidence that IkB α and ϵ preferentially inhibit distinct NFkB family members.⁶⁴ In B cells, IkB ϵ has been shown to play a key role in regulating cRel containing NFkB dimers, with loss of IkB ϵ resulting in increased B-cell survival and proliferation.²¹

Within the TNF pathway, both negative and positive feedback has been reported. Expression of the de-ubiquitinase A20 is strongly NF κ B inducible. However, owing to a long protein half-life and enzymatic effector function, its negative feedback effects do not shape NF κ B dynamics acutely, but rather integrate the history of prior exposure to render the NF κ B pathway less sensitive to subsequent stimuli.⁴² TNF itself is NF κ B inducible, but for full activation, additional signaling mechanisms controlling splicing, mRNA half-life, pro-TNF processing, and secretion must be activated.^{65,66} Hence, TNF functions more like a feed-forward loop in response to PAMPs rather than a positive feedback loop *per se*.

NONCANONICAL NFkB SIGNALING

Signaling via NIK and IKKa

Noncanonical signals are NFkB activating signals that are transduced in a NEMO-independent, but NIK and IKK α -dependent manner. Noncanonical pathways activating signals are primarily developmental signals that activate TNF receptors (BAFFR, CD40, LTβR, RANK, TNFR2, Fn14, etc.), some of which also activate the canonical NFκB pathway.^{67–74} Noncanonical NFkB signaling is known to control a wide variety of developmental phenotypes including B-cell survival and maturation, dendritic cell activation, and bone metabolism.⁷⁵ Several chemokines that regulate lymphoid organogenesis are induced specifically by noncanonical NFkB activation.⁶⁹ Base pair differences in kB sites may contribute to noncanonical pathway-specific gene expression.76,77

While canonical signals transduced by NEMO require phosphorylation-dependent activation of the IKK kinase complex, noncanonical signals transduced by NIK require stabilization and accumulation of the kinase, which, in the absence of signal, is rapidly degraded by a TRAF-cIAP complex.⁷⁸ This ubiquitination-dependent degradation ensures very low basal NIK levels.⁷⁹ Noncanonical stimuli trigger TRAF2-dependent cIAP1-cIAP2 activation, which in turn leads to the proteasomal degradation of TRAF3;

this disrupts the cIAP–TRAF complex reducing NIK degradation and leads to accumulation of NIK.^{79–81} NIK activity is dependent on IKK1, but independent of IKK2. Once activated, the NF κ B-inducing-kinase (NIK) complex has dual roles within the noncanonical pathway (Figure 2(a)). Originally identified as a MAP kinase kinase kinase (encoded by *Map3k14*), NIK activates IKK α by phosphorylation of Ser and Thr residues within an activation loop between subdomains VII and VIII of the kinase domain.^{84,85} Although NIK overexpression results in canonical NF κ B activation,^{85,86} NIK knockouts are not defective in the canonical activation of IKK and NF κ B in response to inflammatory cytokines.⁸²

IκBδ Degradation to Release Preformed Dimers

NIK's first role is in targeting the oligomeric IkB complex, causing IkBo degradation and release of preexisting NFkB dimers for nuclear localization.²⁶ This initial response relies on phosphorylation events and occurs quickly, as IkBo bound to preexisting dimers can release NFkB to localize to the nucleus as soon as it is modified by NIK (Figure 2(b)). While NIK is primarily considered the transducer of noncanonical NFkB signaling and IkBo differs from other IkBs in its ability to inhibit RelB-containing dimers and respond to noncanonical stimuli, IkBô also inhibits RelA-containing dimers. Therefore, NIK-induced IkBo degradation may induce an inflammatory and/or developmental response depending on the existing NFkB dimer repertoire poised within the cell prior to stimulation. Both RelB and RelA induction in response to $LT\beta R$ (noncanonical only) are abrogated in NIK knockout.²⁶

P100 Processing to p52 to Generate New RelB:p52

The second role of NIK is in initiating the proteasome-mediated processing of p100 into p52. Phosphorylation of C-terminal serine residues leads to p100 being recognized by SCF/ β TRCP ubiquitin ligase and subsequent partial degradation of the ARD by the 26S proteasome.^{82,83} This processing produces a mature p52 monomer that is then able to dimerize to form transcriptionally active RelB:p52 and other NF κ B dimers.⁸⁷ It appears that only newly translated p100 is able to be processed into p52,⁸⁸ probably because p100 oliomerizies into a complex that renders it unavailable for processing.²⁶ The multidomain interactions between RelB and p52 (and also p100) co-stabilize both proteins, as RelB protein

(a)

(b)



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FIGURE 2 | The noncanonical nuclear factor κ B (NF κ B) activation pathway. (a) Schematic depiction of the noncanonical NFkB signaling pathway. Developmental signals activate the NIK/IKK1 complex that phosphorylates p100. Most p100 is found in a higher-molecularweight inhibitory complex ($I\kappa B\delta$). Upon phosphorylation, p100 is processed into p52 and is then available to bind RelB, creating a dimer that localizes to the nucleus and binds DNA to activate transcription.^{82,83} Active NIK/IKK1 complex also phosphorylates the p100 within IkBô, resulting in its partial degradation and releasing bound NF_KB dimers for nuclear localization and gene activation.²⁶ (b) Diagram of the regulatory logic of the noncanonical NFκB signaling network. Noncanonical signals activate NIK/IKK1 that suppresses $I \kappa B \delta$ and activates processing of p100 into p52. The suppression of $I\kappa B\delta$ that was sequestering preexisting NF κB dimers results in nuclear localization of NFkB and early-phase gene expression. NIK-dependent p100 processing results in p52 production and the formation of new RelB:p52 dimers that can activate a latephase gene expression response.

levels are decreased in $Nfkb2^{-/-}$ cells and p100 protein is decreased in $RelB^{-/-}$ cells.^{89,90} While p52 is only produced at very low levels in most mammalian

cells, certain cell types, including B cells, show active generation of p52.⁹¹ Processing is tightly controlled by a processing-suppressive region in the C-terminal portion of the p100 protein and disruption of this domain leads to constitutive p100 processing.^{82,92} p100 that does not form p52 is free to form highermolecular-weight inhibitors of NFĸB $(I\kappa B\delta)$. and therefore tight regulation is important to ensure production of IkBo in the absence of noncanonical stimuli to regulate late-phase NFkB activity. NIKinduced p100 processing, and subsequent dimerization with RelB, is a relatively slow process compared to the release of preexisting NFkB dimer from inhibitor, and therefore this results in a late-phase and sustained gene-expression response.

During B-cell maturation, BAFF activates noncanonical NF κ B signaling and the consequence of this is dependent on the state of p100 synthesis within the NF κ B network: at moderate p100 synthesis rates BAFF-induced p52 production fully depletes p100 and prevents the formation of I κ B δ , whereas in the context of high p100 synthesis and resultant I κ B δ formation BAFF causes I κ B δ degradation altering cRel activity and affecting B-cell expansion.⁶¹ Constitutive P100 degradation also contributes to p100 homeostasis and hence is important for a variety of cellular functions.⁹³

ALTERNATIVE NFKB ACTIVATION MECHANISMS

Ribotoxic Stress

Ribotoxic stress, as induced by ultraviolet light (UV) exposure or by unfolded protein response (UPR) inducers, has been found to activate NFkB by inhibiting translation of $I\kappa B\alpha$.^{94,95} Thus, these stimuli engage in crosstalk with inflammatory signaling and amplify NFkB activity (Figure 3(a)). Indeed, UVinduced activation of NFkB was shown to occur in enucleated cells, eliminating UV-induced DNA damage as the primary transducer of this pathway. Instead, in response to UV, eukaryotic initiation factor 2 α (eIF2 α) is phosphorylated through stress response kinases GCN2/PERK. Phosphorylation of eIF2a broadly inhibits transcription initiation, resulting in reduced synthesis of IkBa.94 As free IkBa is constantly degraded and synthesized, the UV-induced reduction in synthesis results in decreased IkBa and amplified NF κ B responses (Figure 3(b)). Interestingly, in response to chronic reactive oxygen species exposure, NFkB may repress prosurvival genes and induce prodeath genes.⁹⁷



FIGURE 3 Nuclear factor κ B (NF κ B) activation by ribotoxic and genotoxic stresses. (a) Schematic depiction of alternative methods of NF κ B activation. Ribotoxic stress inducers lead to the phosphorylation of initiation factor eIF2 α through the action of kinases GCN2 and PERK. Once active, eIF2 α inhibits translation initiation, thereby reducing synthesis of I κ Bs.⁹⁴ The reduction in I κ B leads to more free NF κ B that localizes to the nucleus and binds DNA to promote target gene expression. Genotoxic stress inducers lead to the phosphorylation of ATM and induce complex formation with NEMO in the nucleus.⁹⁶ NEMO is phosphorylated and exported into the cytoplasm where it associates with ELKS and stimulates IKK2-containing complexes. Activation of NEMO/IKK2 complexes results in increased I κ B degradation and localization of NF κ B to the nucleus. (b) Diagram of the regulatory logic of alternative methods of NF κ B activation. UV stress response through GCN2/PERK upregulates eIF2 α which in turn suppresses I κ Bs. The reduced I κ B synthesis reduces the inhibition of NF κ B and increases nuclear NF κ B and target gene expression. In response to DNA damage, ATM is upregulated and activates NEMO through a complex with ELKS. Increased NEMO/IKK2 activation results in suppression of I κ Bs.

Genotoxic Stress

Genotoxic stress also activates NF κ B, although through distinct mechanisms.⁹⁶ In this context, the initiation signal for NF κ B responses originates from within the nucleus and is propagated to the cytosol via mechanisms that involve the nuclear export of upstream signaling molecules. In response to DNA damage NEMO is localized to the nucleus as a result of SUMO-1 attachment.⁹⁸ DNA damage-activated

ATM (ataxia telangiectasia mutated) then phosphorylates NEMO in the nucleus and triggers monoubiquitination of NEMO. ATM then binds to this modified NEMO, which is exported to the cytoplasm and activates IKK to result in degradation of I κ Bs. NF κ B activation upon genotoxic stress is markedly slower and lower in amplitude than that observed for immune receptor signaling, and its physiological role remains unclear.

Shear Stress

Mechanical forces exerted on cells can also activate NF κ B. Shear stress (mechanotransduction) activates NF κ B in osteoblasts through phospholipase pathways that release intracellular Ca²⁺. Activated NF κ B in response to shear stress upregulates COX-2, which plays an important role in the response of bone to mechanical loading.⁹⁹ Hemodynamic forces also exert shear stresses on vascular cells during development of atherosclerosis that result in activation of NF κ B through a mechanism independent from canonical I κ B α degradation.¹⁰⁰

NFKB GENERATION MECHANISMS

NFκB dimeric transcription factors are formed by five monomers (Figure 4). Of the 15 possible dimers, 12 are thought to bind the DNA κB element, and 3 (RelB:RelB, RelB:RelA, and RelB:cRel) form low-affinity intertwined dimers that are unable to bind DNA.¹⁰¹ Of the 12 DNA-binding dimers, 9 contain at least one of the activator proteins, RelA, cRel, or RelB (RelA being the most potent and RelB the least), and generally function as transcriptional activators. The remaining three (the abundant p50:p50 homodimer, and the lesser p52:p52 and p50:p52 homodimers and heterodimers, respectively) may function as activators in conjunction with co-activators, including Bcl3 and IκBζ.

The mechanisms underlying NFkB dimer generation (Figure 4) have only recently received attention, but are critical to understanding how different cell types produce different NFkB dimer repertoires during cell differentiation and development. In some cases, a shift in the dimer repertoire has been documented: for example, B cells shift from a RelA:p50 predominant state at the pre-B stage to a cRel:p50 dominant state in mature B-lymphoid cells and then show strong upregulation of RelB and p52 in terminally differentiated B cells.¹⁰² Also, while most monocyte lineages rely on RelA:p50, GM-CSF-derived inflammatory dendritic cells show high levels of the unusual RelB:p50 dimer, whose generation was shown to be dependent on high constitutive RelB expression and NIK activity.³⁵ These examples show that the NFkB dimer repertoire is responsive to stimuli, albeit at longer timescales (>8 h) than the activation of NFkB from the latent dimer repertoire.

NF κ B family members are obligate dimers; as monomers they are unstable and are thought to be quickly degraded. Thus, the NF κ B dimer generation system is highly dynamic and homeostatic. Below, we summarize the key regulatory mechanisms that



(c) DNA binding and transcriptional activity



FIGURE 4 Mechanisms regulating nuclear factor κ B (NFκB) dimer generation. (a) Diagram of NFkB monomer synthesis and processing.^{10,20} All NF_KB monomers and precursors are NF_KB target genes and induced, to varying extents, by NFkB. RelA/RelB/cRel polypeptides are synthesized in a complete form, ready to dimerize into functional NFkB dimers.^{61,104} p105 is a precursor to p50 that must be cleaved in a process thought to be dependent on IKK2.² p100 must be processed via a NIK/IKK1-dependent pathway into mature p52 before it can dimerize into NF κ B.⁸² (b) Schematic of the NF κ B dimerization process. Monomers must dimerize before they are transcriptionally active. The affinity of binding between monomers varies with two large, activation domain proteins having low affinity. $I\kappa B\beta$ can act as a chaperone, enhancing the effective binding affinity of ReIA to form homodimer by stabilizing this normally weak affinity dimer. (c) Table of the combinatorial composition of potential NFkB dimers, indicating their capacities to bind DNA (indicated by horizontal line) and to activate transcription (indicated by arrows). (d) Diagram of IKK's multiple points of control over NFκB dimer formation. (1) The IKK kinases upregulates monomer expression by activating NF_KB-responsive promoters. (2) IKK1 and IKK2 activities promote processing of p100 to p52 and p105 to p50. (3) IKKs lead to the degradation of IkBs that may function as dimerization chaperones (as for example $I\kappa B\beta$ for ReIA homodimer) as well as inhibitors.

contribute to the NF κ B dimer repertoire. How these function together to determine the specific NF κ B signaling system of specific cell types ought to be a focus of future increasingly quantitative studies.

Dimerization Affinities

Key determinants of the NFkB dimer repertoire are the interaction rate constants between the five monomers. Indirect evidence suggests that the 15 NFkB dimers have dramatically different dimerization affinities, yet remarkably little quantitative information has been published. Recently, analytical ultracentrifugation determined the affinities of the RelA:p50, p50:p50, and RelA:RelA dimers to be in the range of 1-5 nM, 20-50 nM, and 0.8-1.5 µM, respectively.¹⁰³ It is not unreasonable to speculate that other dimers fall into these orders of magnitude, with large dimers (RelA:cRel, cRel:cRel and the DNA-binding incompetent dimers RelB:RelA, RelB; RelB, and RelB:cRel) having low affinities close to the µM range, small dimers (p50:p52, p52:p52) in the high nM range, and dimers composed of one large and one small subunit (RelA:p52, RelB:p50, RelB:p52, cRel:p50, and cRel:p52) having the tightest affinities. However, even if these broad rules prove correct, differences between them are likely to be important in determining the dimer repertoire. Further, it is likely that association and dissociation rate constants, rather than steady-state affinities derived from ratios of these rates, may also be important, as a slow k_{on} rates will lead to a kinetic disadvantage within this potentially competitive dynamical system. If experimental pipelines are established, we expect that posttranslational modifications will likely be found that modulate these important parameters.

Expression of NFkB Monomer Genes

In order to produce NF κ B dimers, a monomer must be expressed, yet given the interdependence of combinatorial dimerization, expression of all monomers must be considered in order to predict the abundance of a single dimer. NF κ B monomer genes are known to be expressed differentially. RelA is thought to be ubiquitously expressed at high levels, whereas cRel is largely restricted to lymphoid lineages (as well as inflammatory dendritic cells), and RelB is also known to be expressed at high levels in specific cell types. All NF κ B genes are to some degree NF κ B inducible, with cRel, RelB, *Nfkb1*, and *Nfkb2* being long-recognized targets of RelA,^{61,104} and RelA recently reported to be a RelA target as well.¹⁰⁵ However, it is unlikely that autoregulation alone accounts for the cell-type-specific expression patterns and little information is available about other transcription factors that control the expression of NF κ B genes.

Processing of NFkB Monomer Precursors

Expression of *Nfkb1* and *Nfkb2* leads to the precursor proteins p105 and p100 that must be processed before the p50 and p52 dimerization partners are available. p105 is endoproteolytically cleaved into a mature protein, p50, that is able to dimerize with other NFkB proteins (predominantly RelA).² p105 processing is generally a constitutive process that occurs in unstimulated cells.¹⁰⁶ It is unclear whether IKK2 may process p105 in a signal-responsive manner.¹⁰⁷

Nfkb2/p100 must be processed into p52 before it can dimerize, predominantly with RelB, to form an NF κ B dimer. In response to developmental signals p52 processing is increased as NIK is activated.⁸² p100 to p52 processing is not only significant as it generates p52 but also as it depletes p100 which would otherwise complex into the inhibitory complex IkBo. Competition for binding to RelA and RelB between p50 and p52 also contributes to p100 processing rate as RelB inhibits p100 to p52 processing.⁹⁰ The level of precursor protein controls the maximal signal-responsive induction of protein processing that can be achieved. When elevated, p100 processing depletes the cellular pool of p100 and noncanonical signaling is unable to strongly induce further p52 production or RelB:p52 formation. Similarly, when all RelB is able to bind to p50 (for example, in $nfkb2^{-/-}$ where no p52 is produced), the pool of precursor p105 is depleted.⁹⁰

Monomer Competition During Dimerization

The combinatorial nature of NF κ B dimerization implies the potential for competition between monomers for the generation of specific dimers. This competition was first reported with regard to the formation of RelA:p50 versus RelB:p52, where a slightly higher affinity of p105/p50 for RelA reduces the opportunity for p100/p52 from complexing with RelA.¹⁰⁸ Conversely, a slightly higher affinity of p100/p52 for RelB diminishes the potential formation of RelA:p52 dimers. This model explains the appearance of such dimers in the respective knockouts.⁹⁰

More recently, a quantitative analysis of RelA homodimerization and heterodimerization (with p50) revealed the degree to which monomer competition

reduces the abundance of low-affinity dimers¹⁰³: the high-affinity RelA:p50 dimer dramatically reduces the abundance of the low-affinity RelA homodimer, whose formation hence becomes entirely dependent on a chaperone. Only when p50 is genetically diminished does RelA homodimer formation become chaperone independent.¹⁰³

Dimer Stabilization and Chaperones

In addition to the combinatorial dimerization affinities and physiochemical properties of the monomers, dimer abundances may be enhanced by 'thirdparty' stabilizers, and dimer generation may be enhanced by dimerization chaperones. To date, one such example has been reported^{103,109}: IkBß was identified as increasing the effective binding affinity of RelA homodimer.¹⁰³ As mentioned, RelA homodimer levels suffer not only from a poor affinity but also from competition from RelA:p50 dimerization. This effect is counteracted by IkBB, which is able to increase the binding of the low-affinity RelA homodimer. A quantitative analysis arrives at such a high effective affinity that a two-step model for IkBß function is plausible, i.e. that $I\kappa B\beta$ binds one monomer first and then a second, enhancing the effective association rate constant and rendering it a bona fide 'chaperone.' However, direct evidence is currently outstanding.

But regardless of whether IkBs function as chaperones or merely stabilizers of specific dimers, their potential role adds a more dynamic and complex component to the control of the cell-type-specific NFkB dimer repertoire, as well as a redefinition of IkBs from being merely inhibitors to also being 'licensing factors' of NFkB activity.

Dimer Degradation

NFκB dimers are thought to be stable when associated with IκBs. They neither come apart nor are they degraded. However, while DNA interactions may also stabilize dimerizing interactions, NFκB dimers bound to DNA are thought to be subject to regulated degradation. Though an attractive hypothesis, the mechanisms remain less than clear. RelA was shown to be removed from specific chromatin sites even in the absence of new IκBα feedback synthesis¹¹⁰; RelA removal from chromatin was impaired in IKKαdeficient macrophages¹¹¹; and the peptidyl-prolyl isomerase Pin1 and the E3 ligases SOCS1¹¹² and COMMD1/Cul2¹¹³ have been implicated in RelA degradation.

SIGNALING CROSSTALK MECHANISMS

While the canonical and noncanonical pathways are mediated by distinct kinases and immediate substrates, given the large number of shared components within the NF κ B system, there is a great potential for crosstalk between the two pathways.

Noncanonical Control of Canonical Signaling

Nfkb2/p100 is the primary signaling node at which canonical and noncanonical signals interact (Figure 5 (a)). That is because (1) Nfkb2 expression is inducible by RelA, (2) any p100 that is not processed into p52 forms a higher-molecular-weight inhibitor of NFκB, the IκBδ-containing IκBsome, that may trap NFkB dimers and thus diminish their association with canonical IkBs, 26,114 and (3) p100 processing to p52 and IkBo degradation is triggered by noncanonical NIK activity. As a result, noncanonical signaling may extend the duration of canonical NFkB activation,⁶¹ or in its absence may diminish canonical NFkB activation.⁵⁸ Thus, noncanonical pathway activity tunes the potency of the canonical pathway in activating NF κ B (Figure 5(b)). Interestingly, in conditions with chronically elevated noncanonical activity, as in inflammatory dendritic cells, canonical pathway signaling is not only altered quantitatively but also qualitatively: in GM-CSF-derived dendritic cells, high noncanonical pathway activity diminishes p100 to such a degree that the RelB:p50 dimer forms, which then associates with $I\kappa B\alpha$ (and to some degree IkBE), rendering it responsive to noncanonical signals.³⁵ Hence, TLR-triggered maturation of these dendritic cells involves activation of RelB:p50, in addition to the expected RelA:p50 and cRel:p50 dimers.

Canonical Control of Noncanonical Signaling

In addition to Nfkb2, the *Relb* gene is also induced by NF κ B dimers in response to canonical pathway activity.^{104,115} Therefore, canonical pathway activity is essential for noncanonical activation of the RelB: p52 dimer.⁹⁰ The dependency of noncanonical signaling on canonical activity impacts lymph node development in RelA-deficient mice.⁸⁷

This potential for crosstalk may in principle also allow elevated canonical pathway activity to amplify noncanonical activation of RelB:p52 (Figure 5(b)), with potentially pathologic



FIGURE 5 Signaling crosstalk between canonical and noncanonical pathways. (a) Diagram of dual roles of Nfkb2/p100 and NIK within the noncanonical pathway that together with the inducible expression of Nfkb2/p100 mediate two crosstalk functions. (1) NIK/IKK1 processes p100 into p52, enabling the activity of RelB.⁸² (2) NIK degrades $I\kappa B\delta$, allowing for sustained RelA activity.⁶¹ (b) Canonical pathway activity may boost noncanonical pathway activation of RelB:p52.⁹⁰ Novel model simulations that illustrate how noncanonical pathway activation of RelB:p52 may be boosted by increasing constitutive canonical pathway activities. (c) A noncanonical pathway stimulus may prolong canonical pathway-induced NF κ B activity. In B cells, BAFF may potentiate late IgM-induced cRel activity.⁶¹

consequences.¹⁰⁸ However, this does not appear to be the case—yet the mechanism that insulates canonical signaling from high canonical activity remains to be elucidated.

suggest a signaling crosstalk that ought to be examined with a range of biochemical, cell biological, and computational research tools.

Posttranslational Modifications

Several posttranslational modifications of RelA have been identified, including phosphorylation of serines and threonines in the DNA binding and activation domains, methylation, acetylation, and glycosylation.^{116,117} Mutational analysis of the modification acceptor residues has in many cases been shown to have detrimental effects on proper NFkB control and target gene expression, and it appears that a variety of regulatory steps (e.g., nuclear localization, DNA binding, and co-activator recruitment) may be affected. However, in many cases, it remains unclear whether the posttranslational modification occurs constitutively, is induced by the same stimulus as IkB degradation, or may in fact be induced by a different stimulus or cell-type specifically. Evidence for the latter scenario would support the attractive hypothesis that NFkB proteins function as integrators of distinct signals (to control nuclear localization and posttranslational modifications) to fine-tune NFkB-responsive transcriptional programs. Such a scenario would

SUMMARY

NFkB has been a key nexus of scientific interest over the past several decades. The breadth and depth of investigation, and the compiled knowledge of this key family of transcription factors are unparalleled. From early insights into its function as a transcription factor and immune response regulator to later studies on cellular signaling, dynamical responses, gene regulatory networks, and feedback mechanisms, it is clear that NFkB has become a standard bearer for research into the inner workings of cellular communication and signal response paradigms. New work on signaling crosstalk and remaining questions and challenges in the field, an amenability to signaling and information transmission studies informed by computational modeling, as well as NFkB's continued interest as a therapeutic target ensure a future of extensive interest and publication on the subject. Ongoing work to expand the scope of computational models and extend their applicability from the cellular scale to the tissue scale will require agent-based

modeling techniques that may also leverage minimal model formulations.^{118–120} Identification and construction of appropriate minimal models for this

purpose depend on identifying the physiologically relevant features within NF κ B's intricate and varied dynamic responses.

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