NO-donating aspirin inhibits the activation of NF-κB in human cancer cell lines and Min mice

Introduction

Cancer prevention emerges as a promising approach to the control of cancer, a major medical challenge of our times. The feasibility and promise of chemoprevention are exemplified by the case of colon cancer. Despite efforts to develop effective therapies, colon cancer, one of the most prevalent cancers in the western world, is the third leading cause of cancer-related deaths in the USA. The limited efficacy of current treatments for advanced colon cancer has provided the impetus for a concerted effort to identify suitable chemopreventive agents. A (partial) list of such agents includes non-steroidal anti-inflammatory drugs (NSAIDs), calcium, antioxidants, polyamine inhibitors, dithiolthiones, polyphenols, vitamin E and selenium (1,2). Epidemiological studies suggested that NSAIDs reduce by about half the incidence of and mortality from colon cancer (3), but recent in-vitro and in-vivo studies of NSAIDs are nitric oxide-donating aspirin (NO-ASA), which is several hundred fold more potent than traditional ASA in inhibiting the growth of colon cancer cells in culture (7). This is also true for cell lines from a variety of tissues such as pancreas, lung, prostate and tongue (8). NO-ASA inhibits the growth of colon cancer cells through a favorable cell kinetic effect that includes decreased proliferation, increased cell death and a block in cell-cycle transitions (9). The mechanism responsible for the enhanced potency of ASA when it is converted to NO-ASA is unknown, but is believed to include effects on cell signaling mediated by cyclooxygenase-2, β-catenin/T-cell factor, nitric oxide synthase-2 and NF-κB (10). Our data on the effect of NO-ASA on NF-κB were of a preliminary nature. Thus, we undertook a detailed study of this potentially significant interaction.

NO-κB is a dimer mostly consisting of the subunits p50 and p65; homodimers of p50, p52 or p65 have also been detected (11). NO-κB is sequestered in the cytoplasm bound to the inhibitory factor IκBz. Phosphorylation of IκBz in response to inducers such as cytokines, reactive oxygen species or infection results in its degradation, unmasking of the nuclear localization signal and activation of NF-κB. As a result, NF-κB translocates to the nucleus and binds to the NF-κB binding site (κB site) in the regulatory region of target genes, thereby promoting the transcription of several regulatory proteins. NF-κB, a strategic signaling molecule in inflammation, promotes cell growth by increasing cell proliferation and inhibiting apoptosis, an effect that directly contributes to carcinogenesis. The nuclear transcriptional factor NF-κB is constitutively expressed in many human cancers, with pancreatic cancer representing a characteristic example (12,13). Over two-thirds of pancreatic cancer display activation of NF-κB, an event considered central to inflammation, a recently appreciated major contributor to carcinogenesis. Of note, NO-ASA is the most effective chemopreventive agent against pancreatic cancer in a preclinical model of pancreatic cancer (14).

Herein, we report our results documenting a profound inhibitory effect of NO-ASA on NF-κB activation, which occurs both in vitro and in vivo and is significantly correlated with NO-ASA’s cell growth inhibitory effect. These findings may explain in part the chemopreventive efficacy of NO-ASA.

Materials and methods

Reagents

We prepared 100-mM stock solutions of para-NO-ASA [2-(acetyloxy)benzoic acid 4-(nitrooxymethyl) phenyl ester] synthesized by us (15) and conventional ASA (Sigma, St Louis, MO) in DMSO; the final DMSO concentration in all experiments of NO-ASA and conventional ASA was 0.1%.

Cell culture

Human colon (HT-29, LoVo, HCT116, HCT-15 and SW-480), breast (MCF-7 and MDA-MB-231) and pancreatic (BxPC-3 and MIA PaCa-2) cell lines (American Type Culture Collection, Manassas, VA) were grown as monolayers in the medium suggested by American Type Culture Collection and supplemented with 10% fetal calf serum (Mediatech, Herndon, VA), penicillin (50U/ml) and streptomycin (50 µg/ml). Life Technologies, Grand Island, NY). Cells were incubated at 37°C in 5% CO₂.

Cell growth

Cell viability/growth was measured using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) colorimetric assay according to the manufacturer’s instructions (Roche Diagnostics, Nutley, NJ). In brief, cells were seeded in 96-well tissue culture plates at a density of 1×10⁴ cells per well and allowed to adhere overnight. On the following day, cells were treated with NO-ASA at concentrations ranging from 1 to 1000 µM for up to 24 h. At the designated time, 10 µl of MTT reagent I was added to the culture media and
the plate returned to the 37°C incubator for 4 h, at which time, 100 µl of the solubilization solution (MTT reagent II) was added and incubation continued overnight. Cellular proliferation was determined spectrophotometrically using an enzyme-linked immunosorbent assay (ELISA) plate reader at 590 nm. The MTT assay was validated in our system by the direct trypan blue method.

**Nuclear extracts**

Nuclear extracts were prepared following standard protocols: trypsinized cells were suspended in lysis buffer to which Nonidet P-40 was added at a subsequent step; nuclei were washed and centrifuged, followed by resuspension in extraction buffer and pelleting (16). Nuclear extracts were stored at −80°C until further analysis.

**Electrophoretic mobility shift assays**

The NF-κB–DNA binding activity was assessed by reacting extracts from control and cells treated with NO-ASA with 33 nmol of a 32P-end labeled 22-mer double-stranded NF-κB consensus oligonucleotide (AGTTGAGGG-GACCTTTCCAGGG; Promega Corp., Madison WI). The specificity of the binding was tested by competition reactions, using 50-fold excess of unlabeled NF-κB (specific) or AP-1 consensus (non-specific) oligonucleotides. The complexes were separated on a 6% gel following standard protocols.

**Enzyme-linked immunosorbent assay**

TransBinding NF-κB assay was performed using an ELISA plate method in which an oligonucleotide harboring a NF-κB consensus binding site was immobilized on a 96-well plate (Panomics, Fremont, CA). Nuclear extracts isolated from colon cancer cells (10 µg) were added to wells of the ELISA plate. The activated NF-κB p50 from the nuclear extract was allowed to bind specifically to the immobilized oligonucleotide. The bound complex was detected by antibody directed against the p50 subunit. A secondary HRP-conjugated antibody was added, followed by addition of the substrate 3,3′,5,5′-tetramethylbenzidine, and the complete complex was quantified by spectrophotometry.

**Western blot analysis**

Proteins were extracted, fractionated by sodium dodecyl sulfate–polyacrylamide gel electrophoresis and transferred onto polyvinylidene difluoride membranes (17). Whole cell lysates were obtained using Radio-Immunoprecipitation Assay buffer (18). Probing with antibodies against the NF-κB subunits p65, p50, IκBα, pIκB-α, Sam68 or β-tubulin (Santa Cruz Biotechnology, Santa Cruz, CA) was done for 1 h at room temperature or overnight at 4°C. The membranes were developed by the enhanced chemiluminescence system (Amersham Biosciences, Piscataway, NJ). Bands were quantified using UVP LABWORKS Imaging Analysis software (UVP LABWORKS, Upland, CA) and normalized to a housekeeping protein.

**Immunofluorescence**

Cells were washed with phosphate-buffered saline and fixed with methanol at 20°C for 10 min. Nuclei were premeabilized by treatment with 0.1% Triton X-100/phosphate-buffered saline for 5 min. Cells were blocked in 2% BSA/phosphate-buffered saline for 15 min and incubated with rabbit p65, p50 or IgG control antibody (Santa Cruz/1:200) for 1 h at room temperature. Subsequently, cells were incubated with Alexa Fluor® 488 anti-rabbit secondary antibody for 1 h at 4°C (1:200). The Zeiss LSM 510 META NLO Two-Photon Laser Scanning Confocal Microscope System was used for detection of staining. The intensity of nuclear staining was determined by using the area measurement function within the histogram tool bar of the Zeiss LSM Image Browser.

**Animal study**

Apcmin/+ mice (Jackson laboratory, Bar Harbor, ME) were treated with NO-ASA (100 mg/kg body weight intra-rectally daily) or vehicle for 3 weeks. Following euthanasia, the colon tissue was immediately preserved in 10% buffered formalin and processed for histopathology.

**Immunohistochemistry**

Tissue samples were isolated, fixed in 10% formalin, embedded in paraffin and sections were placed on slides. The tissue sections were dewaxed and rehydrated according to standard protocols. Tissue sections were pre-treated for antigen retrieval by boiling in a microwave oven for 15 min in a 0.01 M solution of citric acid solution (pH 6.0). Staining was performed according to the instructions for rapid immunohistochemical staining using the VECTASTAIN kit (Vector Laboratories, Burlingame, CA). The primary antibody (Chemicon International, Temecula, CA) detected the activated form of NF-κB, as it specifically recognized the epitope overlapping the nuclear location signal of the NF-κB heterodimer (19). An isotypic antibody control was included with each tissue section.

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Fig. 1. The structure and inhibitory effect of NO-ASA. (A), (B) and (C) Inhibition of NF-κB–DNA binding by NO-ASA. EMSA was performed on nuclear extracts from HT-29 cells treated with NO-ASA for (B) 0.5, (C) 1 and (D) 3 h, as in Materials and methods. (D) Band intensity was determined by densitometry. (E) Inhibition of NF-κB activity by NO-ASA. Nuclear extracts of HT-29 colon cancer cells treated for 3 h with NO-ASA were isolated and the activity of NF-κB was measured by ELISA as in Materials and methods. The IC50 for NF-κB activity as determined under these conditions was 45 µM. This study was repeated twice in duplicate and the results were within 15%. (F) Depiction of the three structural components of NO-ASA (the conventional ASA, -ONO2, the NO-releasing group, and the spacer molecule linking the two). NE, no extract; SC, specific competitor (unlabeled NF-κB oligonucleotide); NSC, non-specific competitor (unlabeled AP-1 oligonucleotide).


Scoring

It was performed using Adobe Photoshop. By using the ‘sampled colors’ function, the positive colors were converted to white and the background was deleted. The histogram function was used to generate the luminosity, which was proportional to the intensity of staining. The gradient luminosities were calculated and used to obtain the ‘mean value’ generating the ‘immunohistochemistry (IHC) luminosity index’. In an independent experiment, we examined the activation of p65 in different tissue sections derived from the same tissue blocks, as described above. The ‘extent of expression’ of activated NF-κB in the epithelial cells of each sample was scored based on the proportion of cells staining positive (0 = no staining, 1+ = up to 1/3 of cells, 2+ = 1/3–2/3, 3+ = 2/3 to 90% and 4+ = 90–100% of cells). We also scored the ‘intensity of staining’ using the following scale: 0 = no staining, 1+ = light brown staining, 2+ = between 2+ and 3+ and 3+ = maximal staining.

Results

NO-ASA inhibits the growth of adenocarcinoma cell lines

In agreement with previous findings, NO-ASA profoundly inhibited cell growth (9). The IC50 for cell growth inhibition in response to NO-ASA, shown in Table I, varied among cell lines originating from a given tissue as well as between cancer types. The IC50 for colon cancer cell lines treated with NO-ASA for 24 h ranged between 10 ± 0.9 and 57 ± 6.3 μM. The IC50s for the pancreatic (range 9.1–22.0 μM) and breast (range 5.2–14.4 μM) cancer cell lines were in general, smaller than those of colon. These results underscore the broad spectrum of the in vitro cell growth inhibitory effect of NO-ASA.

NO-ASA inhibits the activation of NF-κB in various colon cancer cell lines

Given the modulating effect of NF-κB on cell growth, we investigated whether NO-ASA affects the activation of NF-κB in these cell lines. We have observed increase in apoptosis and necrosis in cells treated with NO-ASA for 24–72 h (9). It was, however, important to examine these cells for the effect of NO-ASA on NF-κB prior to the induction of overt cellular damage. Study of NO-ASA induced cell death during incubation periods <24 h revealed that at 3 h there was no apoptosis. Therefore, in most experiments presented here cells were treated with NO-ASA for up to 3 h.

As shown in Figure 1, NO-ASA inhibited NF-κB activation evaluated using electrophoretic mobility shift assays (EMSAs). Such inhibition occurred as early as 30 min and at low a NO-ASA concentration as 10 μM; at 3 h, 50 μM of NO-ASA blocked essentially all NF-κB binding to DNA. The IC50 for this effect was time dependent, being 5.6, 7.8 and 8.6 μM for SW-480, 16.0 ± 2.2 μM for HT-29 and 20.6 ± 1.4 μM for LoVo colon cancer cells, 0.83 μM for HCT-116 and 64 μM for SW-480. Similar to HT-29 cells, the effect of ASA on NF-κB binding was marginal on all these cell lines.

Overall, NO-ASA decreased the binding of NF-κB in a concentration-dependent manner. However, we did observe a slight increase in most cell lines treated with low concentrations of NO-ASA. This increase was consistent between experiments and was not due to unequal sample loading; equal sample loading was confirmed in parallel control samples immunoblotted for the nuclear protein Sam68 (data not shown).

Table I. NO-ASA inhibits the growth of human adenocarcinoma cell lines

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Apc</th>
<th>p53</th>
<th>COX-2</th>
<th>ER</th>
<th>IC50 (μM)</th>
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</thead>
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<tr>
<td>Colon</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SW-480</td>
<td>Mutant</td>
<td>Mutant</td>
<td>–</td>
<td>33.0 ± 5.0</td>
<td></td>
</tr>
<tr>
<td>HT-29</td>
<td>Mutant</td>
<td>Mutant</td>
<td>+</td>
<td>16.0 ± 2.2</td>
<td></td>
</tr>
<tr>
<td>HCT-15</td>
<td>Mutant</td>
<td>Wild-type</td>
<td>–</td>
<td>10.0 ± 0.9</td>
<td></td>
</tr>
<tr>
<td>LoVo</td>
<td>Mutant</td>
<td>Wild-type</td>
<td>–</td>
<td>20.6 ± 1.4</td>
<td></td>
</tr>
<tr>
<td>HCT116</td>
<td>Wild-type</td>
<td>Wild-type</td>
<td>–</td>
<td>57.0 ± 6.3</td>
<td></td>
</tr>
<tr>
<td>Pancreas</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BxPC-3</td>
<td>Mutant</td>
<td>Mutant</td>
<td>+</td>
<td>9.1 ± 0.12</td>
<td></td>
</tr>
<tr>
<td>MIA PaCa-2</td>
<td>Mutant</td>
<td>Mutant</td>
<td>–</td>
<td>22.0 ± 2.0</td>
<td></td>
</tr>
<tr>
<td>Breast</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MCF-7</td>
<td>Wild-type</td>
<td>Wild-type</td>
<td>–</td>
<td>14.4 ± 1.3</td>
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<tr>
<td>MDA-MB-231</td>
<td>Wild-type</td>
<td>Mutant</td>
<td>+</td>
<td>5.2 ± 2.1</td>
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</tr>
</tbody>
</table>

ER, estrogen receptor.

*The expression of COX-2 is barely detectable lasting <30 min.

**The expression of COX-2 is transient lasting <30 min (20) (21).

with NO-ASA for 3 h were quantified for NF-kB activity. Similar to what was observed by EMSA, NO-ASA inhibited NF-kB activity in cells treated for 3 h in a concentration-dependent manner (Figure 1E); the IC50 for this effect was 33.5 μM.

To determine whether the inhibitory effect of NO-ASA on NF-κB binding was unique to HT-29 cells, we examined four additional human colon cancer cell lines differing with respect to their Apc and p53 status and COX-2 expression (Table I). As seen in Figure 2, at 3 h NO-ASA’s IC50 for the binding of NF-κB to DNA was 5 μM for LoVo colon cancer cells, 0.83 μM for HCT-15, 46.3 μM for HCT116 and 64 μM for SW-480. Similar to HT-29 cells, the effect of ASA on NF-κB binding was marginal on all these cell lines.

Overall, NO-ASA decreased the binding of NF-κB in a concentration-dependent manner. However, we did observe a slight increase in most cell lines treated with low concentrations of NO-ASA. This increase was consistent between experiments and was not due to unequal sample loading; equal sample loading was confirmed in parallel control samples immunoblotted for the nuclear protein Sam68 (data not shown).

NO-ASA inhibits the activation of NF-κB in breast and pancreatic cancer cell lines

In much the same fashion as seen with colon cancer cell lines, NO-ASA inhibited nuclear NF-kB binding in breast and pancreatic cells.

As determined by EMSA, NO-ASA decreased the baseline binding of NF-κB to DNA in breast and pancreatic cancer cell lines in a concentration-dependent manner (Figure 3A). In MCF-7 and MDA-MB-231 breast cancer cells, at 3 h NO-ASA decreased NF-κB–DNA binding starting at 10 μM and this inhibition was virtually complete at 50 μM. The IC50s for this effect were 8.9 and 30 μM for MCF-7 and MDA-MB-231, respectively. Similarly, NO-ASA inhibited the DNA binding of NF-κB in the pancreatic cancer cell lines BxPC-3 and MIA PaCa-2 with an IC50 of 11.5 and 7 μM, respectively. These effects mirror NO-ASA’s effect on HT-29 colon cancer cells, shown in Figure 1, suggesting that most probably this effect is independent of the tissue origin of any given cell line.

The role of NF-κB inhibition in cancer cell growth

As shown in Figure 3C, we explored whether the effect of NO-ASA on NF-κB binding to DNA correlated with its effect on cell growth. When we combined all nine cell lines (colon, pancreatic and breast) into one group, the IC50s for NF-κB activation and cell growth were significantly associated (P < 0.05). A similar result was obtained when we restricted our analysis to the five colon cell lines (P < 0.05). These findings suggest that NF-κB probably mediates the cell growth inhibitory effect of NO-ASA on cancer cell lines, regardless of their tissue of origin.

The effect of NO-ASA on the nuclear levels of NF-κB subunits p50 and p65

Supershift analyses demonstrated that in these cell lines the NF-κB heterodimer consists of p50 and p65 (Figure 3B). A non-specific antibody, used as a control, failed to generate the supershift (data not shown). To determine whether the inhibition of NF-κB activation by NO-ASA was due to decreased nuclear amounts of the NF-κB subunits, we monitored the levels of p50 and p65. At NO-ASA concentrations that inhibited the binding of NF-κB to DNA, the p50 and p65 subunits were present at amounts equal to or modestly below control levels (Figure 4A). Therefore, the inhibition of NF-κB binding appears to occur without the loss or appreciable reduction of the NF-κB subunits (p50 and p65) in the nucleus. Results from immunofluorescence analyses of HT-29 and BxPC-3 cells treated with NO-ASA were consistent with those of the immunoblot analyses. As seen in Figure 4B, the fluorescence from p50 and p65 in the nuclei of HT-29 colon cancer cells treated with various concentrations of NO-ASA remained virtually unchanged compared with untreated controls, with the exception of p50 levels that were increased at the highest NO-ASA concentration (100 μM). In contrast, the levels of both p50 and
p65 increased significantly over controls following treatment with TNF-α. Results for the BxPC-3 cell line (data not shown) were similar.

To clarify this unusual situation, we also assayed nuclear levels of IκBα, a protein known to export NF-κB subunits from the nucleus to the cytoplasm by physically associating with them (22,23). As shown in Figure 4A, the nuclear levels of IκBα were increased in response to NO-ASA. In fact, there appears to be roughly an inverse relationship between the inhibition of NF-κB by NO-ASA and the nuclear levels of IκBα. That the nuclear levels of IκBα that we observed are indeed nuclear and not cytoplasmic contaminants generated during its isolation procedure was documented by examining the Sam68 (nuclear) and tubulin (cytoplasmic) control proteins. Thus, it is unlikely that lack of IκBα accounts for maintaining p50 and p65 at around baseline levels. In addition, we examined cellular levels of pIκB. As shown in Figure 4A, the phosphorylation of IκB increased in response to NO-ASA treatment as compared with control levels. Thus, the inhibitory effect of NO-ASA on NF-κB binding cannot be solely attributed to sequestering of NF-κB in the cytoplasm.

We considered that IκBα might engage in a futile physical interaction with the NF-κB subunits such that it prevented their binding to the NF-κB recognition sequence, while also abrogating their translocation into the cytoplasm. Thus, we attempted to communoprecipitate IκBα and p50 or p65 and also to assess their potential colocalization by confocal microscopy. Both efforts proved negative (data not shown).

NO-ASA decreases the activation of NF-κB in Min (Apcmin/+) mice

We assessed whether NO-ASA modulates the NF-κB signaling system in vivo as it does in vitro. Thus, we examined the activation of NF-κB in the colon of 6-week-old female Min (Apcmin/+) mice treated with NO-ASA (100 mg/kg body weight intrarectally daily for 3 weeks). These mice represent an excellent model system of intestinal cancer, as they recapitulate many of its molecular features (24).

Treatment of Min (Apcmin/+) mice with NO-ASA decreased the number of tumors in the small intestine by 59% (P < 0.01), compared with vehicle-treated controls (25). Small intestinal tissue from these animals was examined by IHC, using an antibody that recognizes the activated form of NF-κB. The results are summarized in Figure 5. Compared with controls, NO-ASA decreased the activation of NF-κB by 38.4% (P < 0.01), as determined by the IHC luminosity index. Wild-type control mice treated with vehicle or NO-ASA showed little or no positive staining with this antibody, demonstrating essentially no NF-κB activation. This finding is consistent with the notion that NF-κB is activated during carcinogenesis (26).

In an independent experiment, different sections of the same tissue samples were stained by IHC and scored for NF-κB expression in terms of its intensity and extent, as defined in Materials and methods. NO-ASA decreased the extent of expression of activated NF-κB in murine intestinal epithelial cells by 47% (P < 0.02) and its intensity by 33% (P < 0.05). Both sets of data are in close agreement.

Discussion

Our data indicate that NO-ASA inhibited the activation of N-FκB in a variety of human cancer cell lines through an apparently novel mechanism. Inhibition of NF-κB activation was also noted in an animal model of intestinal cancer.

Our study included human cancer cell lines derived from three clinically important cancers, colon, pancreas and breast, each represented by more than one cell line. In agreement with our previous findings, NO-ASA inhibited potently the growth of all these cell lines (8,10). The presence or absence of COX-2 had no apparent effect on
drug potency, whereas the estrogen receptor positive breast cancer cell line appeared less susceptible (by a factor of three) to the inhibitory effect of NO-ASA compared with the one that was estrogen receptor negative. Conventional ASA, even at millimolar concentrations, had no appreciable effect on cancer cell growth, as noted previously (8,10).

NO-ASA inhibited NF-κB activation as determined either by the EMSA or by an ELISA method assessing its biological activity. Inhibition of NF-κB activation was noted as early as 30 min after exposure to NO-ASA, even before any signs of cell death associated with exposure to NO-ASA. The effect of NO-ASA was clearly concentration dependent as assessed by both EMSA and ELISA. It is of interest, however, that in some cell lines at relatively low concentrations of NO-ASA, NF-κB activation was modestly increased (up to 10–30%) by EMSA. No such stimulation was observed by the ELISA method.

We explored the mechanism of the inhibitory effect of NO-ASA on NF-κB by determining the nuclear levels of its two subunits. Supershift assays revealed that in all cell lines that we tested, the NF-κB dimer was made up of p50 and p65. In the one pancreatic and three colon cell lines that we studied, we demonstrated that over a wide range of NO-ASA concentrations associated with changes in NF-κB activation, the levels of p65 were either unchanged or very modestly decreased compared with controls. A similar observation was made for p50, although at the highest NO-ASA concentrations there was a clear and consistent reduction in its levels. Immunofluorescence analysis of these levels gave similar results, while the positive control (treated with TNF-α) generated the expected rise in p50 and p65 levels. It is clear that for any given concentration of NO-ASA, the NF-κB protein levels in the nucleus cannot account for its reduced binding to the NF-κB DNA recognition sequence. This observation establishes that the reduced NF-κB activity in response to NO-ASA is brought about through a mechanism that does not involve quantitative changes of the NF-κB proteins. In addition, the induction of cytoplasmic levels of pIkB is further evidence that NO-ASA does not reduce the nuclear levels of NF-κB. Thus, the inhibitory binding effect of NO-ASA was not a result of sequestering of the protein in the cytoplasm by IkB. Our conclusion differs from the numerous examples of decreased NF-κB activation owing to diminished nuclear levels of NF-κB subunits [e.g. (27–29)], a mechanism clearly consistent with current understanding of NF-κB regulation.

The possibility that changes in the nuclear levels of IkBα, a protein known to export NF-κB subunits from the nucleus to the cytoplasm (30,31), may account for the observed unusual levels of p50 and p65
in the nucleus is highly unlikely. IκBα levels increased in an apparently inverse proportion to the inhibition of NF-κB by NO-ASA, indicating that p50 and p65 levels were not influenced by quantitative changes in their transport protein IκBα. The possibility that IκBα might prevent NF-κB binding to its recognition sequence while also abrogating the translocation of its subunits into the cytoplasm is also unlikely, as we have failed to demonstrate any such interaction.

We can speculate, for example, that the subunits of NF-κB are modified by NO-ASA in a way that prevents them from binding to DNA. A prime candidate for such a modification is S-nitrosylation of cysteine residues, similar to what we speculated regarding β-catenin (32). Our preliminary data indicate that both p50 and p65 are nitrosylated in response to NO-ASA. In such a case, one could invoke steric hindrance from cysteine S-NOs affecting the κB-binding domain of the protein. Alternatively, conformational changes of one or both NF-κB subunits as a result of S-nitrosylation could explain the observed results.

NF-κB is known to modulate cell proliferation and cell death, especially by apoptosis, depending on biological context (e.g. cell or tissue type) (26). Our previous work with these and other cell lines (33) has demonstrated that NO-ASA inhibits cell growth through its combined effects on cell renewal and cell death. Our demonstration that the IC50 for inhibition of NF-κB is significantly correlated with the IC50 for the inhibition of cell growth is in keeping with such a role of NF-κB in cell kinetics. This association was statistically significant when we combined all nine cell lines (colon, pancreatic and breast) into one group and when we examined the five colon cell lines as a separate group. This observation indicates that the association between NF-κB and cancer cell growth in response to NO-ASA is probably a generalized property.

Based on our data, it is difficult to conclude that the effect of NO-ASA on NF-κB is responsible, alone or in combination with other effects, for its pharmacological effects against cancer. Nevertheless, such a possibility is plausible. NF-κB is an important factor in carcinogenesis through both its modulation of cell kinetics and its central role in inflammation; NF-κB is currently being appreciated as an important link between inflammation and cancer (34). For example,
NF-κB is activated in over two-thirds of human pancreatic cancers, participates in early events of pancreatic carcinogenesis through its interactions with signaling pathways and suppression of its activation restores pancreatic cell kinetics, mainly normalizing the suppressed apoptosis of pancreatic cancer (reviewed in ref. 35).

In conclusion, our findings document a quantitatively significant in vitro inhibitory effect of NO-ASA on the activation of NF-κB in a variety of cell lines from three important human cancers, colon, pancreas and breast. This effect was also observed in vivo in a model of intestinal cancer. Although the mechanism of this inhibition is not yet deciphered, it is clearly different from that involving reduced nuclear levels of the constituent proteins of NF-κB. Since NF-κB inhibition by NO-ASA correlates with its cell growth inhibitory effect, this effect may have direct relevance to the chemopreventive effect of NO-ASA and, at least in part, may explain it.

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References


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