Enzymatic mechanism of the tumoricidal action of 4-iodo-3-nitrobenzamide

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Abstract. Activation of the prodrug 4-iodo-3-nitrobenzamide critically depends on the cellular reducing system specific to cancer cells. In non-malignant cells, reduction of this prodrug to the non-toxic amine occurs by the flavoprotein of complex I of mitochondria receiving Mg$^{2+}$-ATP-dependent reducing equivalents from NADH to NADPH via pyridine nucleotide transhydrogenation. This hydride transfer is deficient in malignant cells; therefore, the lethal synthesis of 4-iodo-3-nitrosobenzamide takes place selectively. Enzymatic evidence for this mechanism has been provided by cellular studies with lyssolecithin-permeabilized cells and cell fractions, which have identified the defect in transhydrogenation in neoplastic cells to be located at the energy transfer site. Confirming previous results, the present study demonstrates the validity of the selective tumoricidal action of the prodrug in cell cultures.

Introduction

Identification of the chemical and enzymatic (by cytochrome P-450) oxidation of Aryl amines to their first oxidized product, nitroso Aryls (1), marked a significant milestone in our investigations aimed at the chemotherapy of neoplastic diseases. We showed that C-nitroso Aryls induced precipitous cell death in tumor cells (2), an effect which was so rapid that tumor cells exposed to these drugs autolysed overnight in culture dishes. However, the cell killing by synthetic nitroso Aryls of neoplastic cells was only 2- to 3-fold greater than that observed in non-malignant cells, therefore exhibiting in vivo toxicity. For these reasons, we developed non-toxic nitroso precursors that are enzymatically reduced to the cytotoxic nitroso derivative in neoplastic cells only. A prototype nitroso precursor is 4-iodo-3-nitrobenzamide (4-I-3NO$_2$BA) (3,4). It must be emphasized that the vicinal iodo-nitro substitution in 4-I-3NO$_2$BA eliminates a direct poly(ADP-ribose) polymerase (PARP-1, E.C. 2.4.2.30) inhibitory action of this benzamide derivative. PARP-1 inhibitory propensity is common to numerous benzoamides, which compete with the nicotinamide moiety of NAD$^+$ (5). The acid-amide group in 4-I-3NO$_2$BA enhances rapid cellular penetration over the carboxylate (3,4), which is the reason for retaining the acid-amide group.

The fate of the cell type exposed to 4-I-3NO$_2$BA depends on its xenobiotic metabolism: cancer cells are killed while non-malignant cells experience no harmful consequences. The predictable xenobiotic metabolism of 4-I-3NO$_2$BA is its reduction to the corresponding non-toxic 4-I-3NH$_2$BA, which is its reduc tion. PARP-1 inhibitory propensity is common to numerous benzoamides, which compete with the nicotinamide moiety of NAD$^+$ (5). The acid-amide group in 4-I-3NO$_2$BA enhances rapid cellular penetration over the carboxylate (3,4), which is the reason for retaining the acid-amide group.

Since the enzymatic reduction of R-NO$_2$ to R-NH$_2$ hinges on the catalytic activity of the multifunctional mitochondrial flavoprotein of complex I. The energy coupled reduction of this flavoprotein by NADPH depends on the transhydrogenation of NADP$^+$ by NADH (8). Since we observed an as yet unspecified bioenergetic defect in cancer cells (9), we predicted that this defect may impede the OXPHOS-dependent reduction of R-NO$_2$ to R-NH$_2$ in tumor cells. Energy-linked pyridine nucleotide transhydrogenation has been extensively studied in mitochondria isolated by the sucrose gradient centrifugation technique and in mitochondrial fragments and purified systems. However, the sonication-dependent isolation technique of submitochondrial fragments (8) failed to separate soluble and particle-bound components of the transhydrogenase and produced truncated fragments. On the other hand, treatment of intact cells with lyssolecithin (9,10) allowed us to study the regulation of transhydrogenation in nearly intact cells, a feat which has not been accomplished thus far, allowing us to pinpoint the difference in transhydrogenation between malignant and non-malignant cells at the particulate fraction, containing components of energy transduction. These results confirm the findings of previous studies (9), which showed defective bioenergetics in malignant cells in cell cultures and in vivo.

Since the enzymatic reduction of R-NO$_2$ occurs through the flavoprotein of complex I, which receives reducing equivalents by way of the energy-dependent formation of NADPH from NADH, we concluded that the metabolic specificity of tumor cell killing by 4-I-3NO$_2$BA is directly attributable.

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to its reduction to 4-I-3NOBA. The present study provides experimental evidence to support this enzymatic mechanism.

**Materials and methods**

**Synthesis and chemical analyses.** The synthesis of 4-I-3-NO$_2$BA and chemical methods of analyses were identical to those previously reported (3,4,6,7). Lysolecithin (LL) was purchased from Sigma-Aldrich, and 3-acetylpyridine adenine dinucleotide (reduced form) (AcNADH) was from OYC Americas (Andover, MA, USA).

**Cell culture.** Cell culture methods for CV-1 (9,10) and Eras 20 cells were identical to those previously reported (11). All cancer cells, upon prolonged subculturing, undergo varying degrees of senescence. This modifies their malignant phenotype (12). In contrast, Eras 20 cells obtained by the transformation of bovine aortic endothelial cells (11) for 25 years showed no signs of senescence, and for this reason were used as an invariable neoplastic prototype cell.

**Permeabilized cells and cell fractions.** Permeabilized cells and cell fractions were prepared as previously reported (9,10). Briefly, CV-1 and Eras 20 cells were detached and counted from appropriate cultures, and cells were suspended in a standard buffer consisting of 150 mM sucrose, 80 mM KCl and 35 mM HEPES buffer of pH 7.4. The cell number/buffer volume ratio was 10$^6$ cells/100 µl buffer, to which 0.21 mg/ml LL was added from an LL stock solution (1 mg/ml). Cell suspensions were kept on ice (0–4°C) during this manipulation. Cell fractions [soluble SU and particulate (M$x$)] (10) were prepared in Eppendorf centrifuge tubes by sedimentation (Eppendorf model 5415R centrifuge at 4°C at 6600 rpm for 6 min). After the quantitative removal of the SU fraction, the M$x$ sediment was resuspended in the LL standard buffer (see above) at a density of 10$^6$ cell equivalents/100 µl. This technique enabled us to dispense 10$^6$ cell equivalents for each spectrophotometric enzyme assay.

**Spectrophotometric transhydrogenase tests.** Two assay systems were developed. The first contained 0.2 mM AcNADH, 0.38 mM NADP$^+$, 1 mM Mg$^{2+}$-ATP, 10 mM deoxyglucose (9) and LL-cells (or cell fractions) equivalent to 10$^6$ cells in Tris-HCl (25 mM, pH 7.5) in a volume of 1.0 ml (Fig. 1A). Mg$^{2+}$-ATP was an equimolar mixture of 1 mM MgCl$_2$ and 1 mM ATP. The Mg$^{2+}$ complex of ATP was an absolute requirement for the transhydrogenation of AcNADH to NADP$^+$; ATP alone was ineffective. Tests were performed in a Shimadzu UV-1650 PC system (10) in 1 ml quartz cuvettes of 1 cm light path and 1 ml volume at 25°C. The rates of NADPH formation were followed at 340 nm, and the generation of AcNADH at 268 nm.

In the second assay system, AcNADH was omitted and replaced by cellular NAD$^+$ and reducing substrates, both present in LL-cell suspensions (Fig. 1B), thus representing a metabolically generated reducing equivalent producing system. The comparison of two transhydrogenation assays in whole cells appeared necessary to establish that no rate limiting factors in the whole cells interfered with the $V_{max}$ of hydride transfer, and the assays actually measured only the rates of transhydrogenation. The combination of both assays makes this method applicable to any cell type.

**Quantitative assays.** Quantitative assays for the cytocidal action of 4-I-3NO$_2$BA were carried out as previously reported (3,4).

**Results and Discussion**

Rates of NADPH formation by LL permeabilized cells determined by added AcNADH (Fig. 1A) or metabolically generated cellular NADPH (Fig. 1B) are presented in Fig. 1. It is apparent that both test systems yielded nearly identical $V_{max}$ rates (n=4, SD ±8%) when suspensions of non-malignant CV-1 cells were compared. The agreement of the two transhydrogenase tests indicates that the generation of intracellular NADH as a precursor of NADPH or the availability of AcNADH is not rate limiting in transhydrogenation. The exact enzymatic reaction mechanism that yielded metabolically-generated cellular NADH capable of reducing NAD$^+$ is not known and is the subject of further investigation, particularly since the NAD$^+$-reducing metabolite may vary in various cell types and could contribute to cellular metabolic specificity.

According to the perspective of current studies, it is evident in Fig. 1 that Eras 20 cells exhibit only marginal Mg$^{2+}$-ATP-dependent NADH-NADP$^+$ transhydrogenase activity.

The nature of this defective transhydrogenation in Eras 20 cells is illustrated in Fig. 2A and B. As we previously reported (9,10), rapid centrifugation of an LL-permeabilized cell suspension separates a particulate (M$x$) fraction containing nuclei and perinuclearly located mitochondria and a soluble SU fraction (10). The M$x$ fraction, which retains subcellular structures noted by electron microscopy (9) and the soluble SU fraction, both prepared from CV-1 cells, each had marginal energy-dependent transhydrogenase activity when tested separately (Fig. 2A). However, their recombination yielded maximal enzymatic rates (Fig. 2A), demonstrating that the LL-fractionation technique of CV-1 cells retained protein sites in both fractions. Upon recombination, these yielded full transhydrogenase activity to NADP$^+$ present in unfractionated permeabilized cells.

Fig. 2B illustrates the results of enzymatic tests conducted with cell fractions prepared from Eras 20 cells combined with those prepared from CV-1 cells. The SU fraction of CV-1 cells yielded only 30% of transhydrogenase activity when combined with the M$x$ fraction of Eras 20 cells (lower curve of Fig. 2B), while the SU fraction of Eras 20 cells when combined with the M$x$ fraction of CV-1 cells recovered full transhydrogenase activity. As a complement to these tests, we determined the ‘energy-independent’ H transfer from NADPH to AcNAD$^+$ assayed as the reduction of AcNAD$^+$ by NADPH at 268 nm, and found them to be equally active in the SU fractions of CV-1 and Eras 20 cells (data not shown). The sum of these results demonstrates that the pyridine nucleotide transhydrogenase enzyme in the SU fractions of CV-1 and Eras 20 cells was equal. Consequently, depressed Mg$^{2+}$-ATP-dependent transhydrogenase activity in combined SU fractions from CV-1 cells and M$x$ fractions from Eras 20 cells (Fig. 2B) clearly indicates that the major enzymatic defect is in the M$x$ fraction of Eras 20 cells. This defect results in an inhibition
of approximately 70% of transhydrogenation (compare curves in Fig. 2B). Loss of 70% activity is apparently enough for the accumulation of the cytocidal nitroso species in tumor cells.

It is of interest that the requirement of inhibition of cellular respiration by CN⁻ as reported by some investigators (8) was unnecessary for the assay of Mg²⁺-aTP-dependent transhydrogenation in the LL-cellular system (Figs. 1 and 2), indicating that the increased maintenance of cell structures favors the effectivity of Mg²⁺-aTP.

The requirement of Mg²⁺-aTP for the NADH→NADP⁺ transhydrogenase reaction in LL cells poses some unresolved questions in cellular bioenergetics, since it is unclear why external Mg²⁺-aTP is necessary in cells where OXPHOS is intact (e.g., in CV-1 cells). Externally added Mg²⁺-aTP to this system appears to imply a biochemical link between mitochondrial ATP synthase and extramitochondrial Mg²⁺-aTP, where ATP is most likely to be derived from glycolytic ATP synthesis. This is a problem in cellular bioenergetics that is the subject of our further studies. However, the unavailability of details regarding the interplay of mitochondrial and extramitochondrial ATP generating systems does not interfere with the interpretation of the mechanisms of the lethal synthesis of iodo-nitrosobenzamide from 4-i-3NO₂Ba prodrug in cancer cells.

This tumoricidal mechanism involves first the generation of the nitrosylating species (3,4), and also that of GS-sulfinic acid without deiodination. GS-sulfinic acid was recently shown to be the most potent inhibitor of the ubiquitous gSH-transferase (13). This transferase can act on nitrosylated targets of the prodrug, thereby eliminating cytocidal effectivity. Inhibition of the transferase therefore potentiates the action of the prodrug.
specific metabolism has led to the discovery of the new mode of chemotherapy described in this study, illustrating the promising future of this aspect of cancer pathology.

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References


