COMPARATIVE KILLING EFFICIENCIES FOR DECAYS OF TRITIATED COMPOUNDS INCORPORATED INTO E. COLI

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ABSTRACT The killing efficiencies due to the decay of incorporated H^* -thymidine, H^* -uridine, and H^* -histidine in E. coli 15_{T-L} have been determined. Decays from H^* -thymidine are 2.0 times as effective in producing lethality as those from H^* -uridine and 2.5 times as effective as those from H^* -histidine. Therefore, it seems that the greater part of damage from H^* -thymidine decays is due to chemical changes associated with nuclear transmutation.

INTRODUCTION

It has been shown that decays from incorporated H³-thymidine in $E.\ coli\ 15$ auxotrophs cause death at a rate approaching that due to P³² decay (Person and Lewis, 1962). A similar result has been reported by Cairns (1961) for H³-thymidine decays in bacteriophage, while Apelgot and Latarjet (1962) have reported a smaller rate of killing in bacteria. Although direct evidence is lacking, the relatively large killing efficiency for decays from H³-thymidine suggests that lethality may be due to other factors in addition to radiation damage. If lethality is not caused exclusively by β -particle ionizations that accompany the nuclear transmutations it may be due to a chemical change in the thymidine molecule. The recoil He nucleus would not be suspect because it is biologically unimportant, chemically unreactive, and has an average energy of about 1 ev, which is probably too low to rupture covalent bonds.

It is the purpose of this communication to describe experiments which were designed to determine the extent of radiation damage due to decays from H⁸-thymidine. Three H⁸ compounds were used so the label would be incorporated into the bacterial DNA (H⁸-thymidine), RNA (H⁸-uridine) or protein (H⁸-histidine). The highly radioactive bacteria were stored, and the efficiency of killing, the probability that a single radioactive decay will produce a lethal event, was determined for H⁸ decay from each of these H³ compounds.

If the major source of lethality is due to radiation damage we postulated that the killing efficiency, which is on a per decay basis, would be nearly the same for each of the three H³ compounds. If factors such as chemical rearrangements are important we would expect a different killing efficiency for each H³ compound.

We have found that decays from H³-thymidine are 2.0 times as effective in producing lethality as those from H³-uridine and 2.5 times as effective as those from H³-histidine.

MATERIALS AND METHODS

A mutant of strain 15, 15_{T-L} , which is thymidine- and leucine-deficient, was used in these experiments. Cultures of this mutant were obtained from Dr. F. Forro, Jr., Yale University. Stock cultures were grown on A-1 medium: 2 gm NH₄C1, 6 gm anhydrous Na₂HPO₄, 5 gm NaCl, 0.115 gm Na₂SO₄, 0.34 gm MgCl₂.6 H₂O, 4 gm glucose, 1 litter H₂O. Experimental cultures were supplemented with leucine at 40 μ g/ml and with either thymidine or H³-thymidine at 2 μ g/ml.

The labeling procedure and the procedure for determining the number of H^a atoms/bacterium have been described previously (Person and Lewis, 1962). Briefly, they are as follows: The desired quantity of a H^a compound was added to a log phase culture growing at 37°C with aeration. The culture was grown in the presence of the radioisotope for 4 to 5 cell divisions. At this time the remaining exogenous H^a compound was separated from the cells and the cells stored at H^a C in A-1. Cell counts and H^a counts were made directly on this highly radioactive culture from which H^a , the number of H^a atoms/bacterium can be computed. Dilutions of stored cultures were made at various times in A-1 medium and platings for viability were on both A-1 supplemented and nutrient agar.

Cells labeled with H³-thymidine or H³-uridine were extracted with trichloroacetic acid (TCA) and NaOH to separate their nucleic acids. This procedure, due to Schmidt and Thannhauser (1945), involved 0.3 M TCA extraction at 4°C (acid-soluble fraction), overnight incubation at 37°C in 1 N NaOH (RNA degradation), heating in boiling water bath for 30 minutes in 0.3 M TCA (DNA degradation), and the uptake of the remaining residue in 1 N NaOH. Enzymatic digestion with 100 μ g/ml of RNAase and/or 50 μ g/ml of DNAase at 37°C for 30 minutes also used in conjunction with cold TCA extraction.

RESULTS

Survival curves for H³ decay in bacteria are "single hit" and satisfy the relation $S/S_0 = \exp(-\alpha_{H^3} \lambda N^* t)$ or $\ln S/S_0 = -\alpha_{H^3} \lambda N^* t$ where S is the number of cells viable at time t and S_0 is the number originally present at t = 0. α_{H^3} is the probability of a lethal event being produced by a single H³ decay. Since α_{H^3} reflects the

¹ H⁸-thymidine, 3.0 c/mmole; H⁸-uridine, 0.9 c/mmole; and H⁸-histidine, 1.7 c/mmole were all obtained from Schwarz Bioresearch, Orangeburg, New York. H⁸-thymidine and H⁸-histidine were used at the above specific activities, but H⁸-uridine was diluted to a specific activity of 0.4 c/mmole. Final concentrations were H⁸-thymidine, 2 μ g/ml; H⁸-urdine, 28 μ g/ml; H⁸-histidine, 4.5 μ g/ml.

lethal yield per decay it has come to be known as the killing efficiency. λ is the decay constant for H³, and N* is the number of H³ atoms/bacterium at time t. However, N* is constant with time in these experiments because $\lambda = 1.52 \times 10^{-4}$ /day and the experiments last only about 8 days. The value of N* is determined by measuring the cells/ml and the H³/ml for an aliquot of stored radioactive cells.

When λ is expressed as the fraction of radioactive atoms that decay per hour, λN^* is the number of decays/hour/bacterium; so that λN^*t is the number of decays/bacterium at any time t in hours. Therefore, if we plot the fraction of cells surviving on a logarithmic scale (S/S_0) as a function of decays/cell (λN^*t) , a straight line results whose slope is α_{H^3} . α_{H^3} may be calculated directly since the assay for viability measures S/S_0 as a function of time; λ is a constant, and N^* is a constant for any one experiment. When the data are plotted in this way differences in the slopes of survival curves reflect different killing efficiencies. This procedure was used to determine α_{H^3} for the three H³-labeled compounds.

Fig. 1 is a semilogarithmic plot of the surviving fraction of $15_{\text{T-L-}}(S/S_0)$, labeled

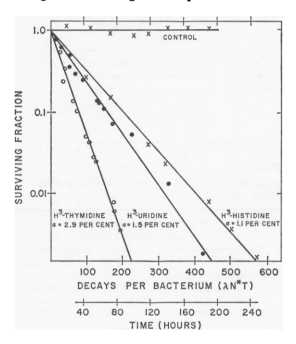


FIGURE 1 The survival of $15_{T.L.}$ as a function of exposure to decays from H³-thymidine, H³-uridine, or H²-histidine. After growth in medium containing one of these H³ compounds the bacteria are stored at 4°C in A-1. Since the exposure to decay is plotted as decays/bacterium ($\lambda N*t$) the differences in slopes of the survival curves reflect different killing efficiencies. It is clear that the killing efficiency for H³-thymidine decays is markedly greater than for that from either H³-uridine or H³-histidine.

with H⁸-thymidine, H⁸-uridine or H³-histidine, as a function of the number of decays/bacterium (λN^*t). After growth in medium containing one of these compounds the bacteria are stored in A-1 at 4°C without supplements. Data showing the viability of an unlabeled culture over the same time period are also shown in Fig. 1. The average killing efficiencies as well as typical values for H³ atoms/bacterium and decays/hour/bacterium are listed below.

Quantity	H³-thymidine	H ⁸ -uridine	H ⁸ -histidine
N*(H* atoms/bacterium)	3.3 × 10 ⁵	2.4 × 10 ⁵	1.05 × 10 ⁶
λN*(H² decays/hr./bacterium)	2.1	1.5	6.7
α _H s(per cent)	2.9	1.5	1.1

The probability of producing lethality per decay, under these conditions, is 2.9 per cent for decays from H⁸-thymidine, 1.5 per cent for decays from H⁸-uridine, and 1.1 per cent for decays from H⁸-histidine.

The results of cell fractionation studies are shown in Tables I and II. Radioactivity determinations were made in a liquid scintillation counter.

TABLE I
THE DISTRIBUTION OF HATHYMIDINE AND
HAURIDINE IN E. COLI 15_{T-L-}
USING THE SCHMIDT-THANNHAUSER PROCEDURE

	Amount of H³-labeled compound, per cent of total activity		
Treatment and fraction	H ⁸ -thymidine	H ⁸ -uridine	
Soluble in cold TCA (acid-soluble)	5.6	6.5	
Soluble in warm NaOH (RNA)	1.7	84.2	
Soluble in hot TCA (DNA)	85.0	8.8	
Residual fraction	7.8	0.6	

TABLE II

THE DISTRIBUTION OF HATHYMIDINE, HAURIDINE, AND HAHISTIDINE IN E. COLI 15_{T-L-} USING ENZYMATIC DIGESTION

	Amount of H³-labeled compound, per cent of total activity			
Treatment and fraction	H ³ -thymidine	H³-uridine	H³-histidine	
Soluble in cold TCA (acid-soluble)	1.7	9.9	10.9	
Soluble in RNAase (RNA)	5.4	88.3		
Soluble in DNAase (DNA)	90.9	1.7	_	
Soluble in RNAase and DNAase (DNA + RNA)	_		1.9	
Residual fraction (protein)	1.9	0.1	87.8	

DISCUSSION

The data presented above show that decays from H³-thymidine are more effective in producing lethality than those from either H³-uridine or H³-histidine. Since the killing efficiencies for the different H⁸ compounds are not the same we feel that

radiation damage is not the sole cause of killing for the H3-thymidine decays. If we assume that the smallest α_{H^3} , for H³-histidine decays, is entirely due to radiation damage, then we can calculate the fraction of killing for H3-thymidine and H3uridine decays that is not due to radiation damage. Since α_{H^3} (thymidine) = 2.9 per cent, $\alpha_{\rm H^3}$ (uridine) = 1.5 per cent, and $\alpha_{\rm H^3}$ (histidine) = 1.1 per cent; 1.8/2.9 of the H3-thymidine-induced damage and 0.4/1.5 for H3-uridine-induced damage would have a cause other than radiation. On the other hand, radiation damage would account for 1.1/2.9 for H³-thymidine- and 1.1/1.5 for H³-uridine-induced lethality. It would seem that about 3/3 of the damage from H3-thymidine decays is not radiation-induced, but is more likely due to chemical changes in the thymidine molecules. H3-thymidine decays are probably more efficient than those from H⁸-uridine or H⁸-histidine because of their strategic location in the "genetic part" of the cell's DNA. While in all cases approximately 85 per cent of the H3labeled compounds used are incorporated into the expected macromolecular fractions, it is possible that a small fraction of damage from H⁸-uridine decays is due to a chemical alteration of the uridine molecules following H³ decay. If true, this could be due to the conversion of a small amount of H³-uridine to H³-deoxycytidine with its subsequent incorporation into DNA. Caro and Forro (1961), using 15_{T-U}. reported that 8 per cent of H8-uridine was incorporated into DNA, and all of this was in the cytosine fraction. Our data are in agreement with this figure.

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