diphenyl-3,3'-diindolymethane. Dahlbom and Misiorny were quick to reply to our unworthy suggestion, and in the light of further work to be described here, we offer our sincere apologies to these authors for suggesting that they had failed to perform a mixed melting point between 2-phenylindole and the product, m.p. 184–185°.

In the present work 2-phenylindole was refluxed in ethanol, containing a trace of hydrochloric acid, with ethyl orthoformate for 24 h. The product was 2,2'-diphenyl-3,3'-diindolymethane which was isolated as its perchlorate (II) according to the procedure of Harley-Mason and Bu'Lock. Crystallization from acetic acid afforded orange-red prisms, m.p. 289–290° (decomp.). (Found: C 69.83; H 4.34; N 5.58. Calc. for C₂₅H₁₈O₅N₂: C 70.09; H 4.26; N 5.64.) This methene salt was hydrolyzed in ethanol in the presence of platinum oxide. Evaporation of the filtered solution yielded I which was obtained as colorless needles from ethanol, m.p. 188–189° (Found: C 87.50; H 5.77; N 6.78. Calc. for C₂₇H₂₈N₂: C 87.46; H 5.75; N 6.75). The di-1,3,5-trinitrobenzene complex of 2,2'-diphenyl-3,3'-diindolymethane was obtained as reddish brown needles from methanol, m.p. 161–162° (Found: C 59.86; H 3.62; N 13.82. Calc. for C₂₅H₁₈O₅N₂: C 59.71; H 3.42; N 13.59). In order to confirm that we had obtained authentic I, the compound m.p. 188–189° was N-methylated according to the procedure of Potts and Saxton to yield 1,1'-dimethyl-2,2'-diphenyl-3,3'-diindolymethane, m.p. 185–186°, identical (mixed m.p., I.R. spectrum) with material previously obtained by the self-condensation of 3-hydroxymethyl-1-methyl-2-phenylindole.

It is evident that Dahlbom and Misiorny obtained authentic I (the slight difference in our observed melting points for this compound is probably not significant) and it is remarkable that 2-phenylindole, 2,2'-diphenyl-3,3'-diindolymethane and its 1,1'-dimethyl derivative should melt so close together. We have also investigated the self-condensation of 3-hydroxymethyl-2-phenylindole. In agreement with our previous results we obtained 2-phenylindole when this compound was refluxed with 10% sodium hydroxide. However, when exactly neutral or slightly acidic conditions were used the product was in fact the diindolymethane I. We suspect that traces of alkali caused the formation of 2-phenylindole when 3-hydroxy-

methyl-2-phenylindole was refluxed with water in our previous experiments.

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Acid Soluble Nucleotide Linked Peptides in Extracts of Ascites Tumor Cells

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In a recent paper it was reported that trichloroacetic acid (TCA) extracts of rabbit livers contained at least four ultra violet (UV) absorbing peaks with bound ninhydrin positive material. These four fractions were eluted immediately after adenosine diphosphate (ADP) and guanosine diphosphate (GDP) with a formic acid-ammonium formate system from a Dowex I column. The positions on the elution curve of the UV-linked ninhydrin positive material were in the same region where we previously had found the uridine nucleotide bound peptides in L. casei. Using a modification of the method of Strominger and Thrann it was possible to isolate from the four fractions ten UV-absorbing spots. Seven of these contained from three up to seven amino acids, some of them at least bound to uridylic acid.

Since these data seemed to prove that animal tissues also contained acid soluble nucleotide bound peptides of a type previously only observed in extracts of different

Fig. 1. Acid soluble nucleotides from $3.8 \times 10^9$ tetraploid tumor ascites cells. The nucleotides were separated on a 3 x 50 cm Dowex 1 formate column by gradient elution with reservoir content changed at the tubes numbered as follows: 72, 1 N formic acid; 164, 4 N formic acid; 390, 0.2 M ammonium formate + 4 N formic acid; 510, 0.4 M ammonium formate + 4 N formic acid. The continuous line represents radioactivity and the broken line represents optical density values at 260 nm ($E_{260}$).

microorganisms it was of interest to look for them in other animal cells. In connection with other experiments the TCA-extracts of ascites tumor cells were chromatographed in the usual way. In Fig. 1 a section of an elution diagram is presented consisting of the peaks immediately before and after ADP. The tumor cells had been separated by counter-streaming centrifugation (by prof. Lindahl) into mainly tetraploid ($3.8 \times 10^9$) and octoploid cells ($10.9 \times 10^9$) and incubated for 30 min. with $^{32}P$ in relation to cell numbers. The TCA extract in Fig. 1 was from the tetraploid cells. The TCA extract from the octoploid cells qualitatively gave a similar elution diagram. In view of the low UV-absorption values the X_{III} and X_{III} peaks from the tetraploid cells were taken together with the corresponding peaks from the octoploid cells. After charcoal-absorption and elution less than 0.5 mg material was obtained from each of the combined X_{III} and X_{III} peaks. No attempt was therefore made to separate the material on an electrophero-chromatogram. After acid hydrolysis all material was put on each of two papers for two dimensional paper chromatography. Both papers showed stronger spots in positions corresponding to glycine, serine, glutamic acid and aspartic acid. Weaker spots were obtained in the positions of leucine or isoleucine, lysine, arginine and threonine. At least four more unidentified spots were also observed on both papers.

Preliminary estimations of the specific activity calculated as counts per minute per \mu mole of nucleotide compound gave the following values:

<table>
<thead>
<tr>
<th>Type</th>
<th>Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tetraploid</td>
<td>$40 \times 10^4$</td>
</tr>
<tr>
<td>Octoploid</td>
<td>$11 \times 10^4$</td>
</tr>
</tbody>
</table>

The figures are of interest since the tetraploid cells are the more rapidly dividing of the two cell types and nucleotide-linked peptides in microorganisms have been claimed as precursors in the formation of cell wall proteins.

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7. Lindahl, P. E. *Personal communication*.

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