in one single operation by means of gradient elution. If the analysis of the eluate is performed spectrophotometrically at 325 mμ, where the contributions from other hop substances is small, a solvent extract of hops can be used without purification. Fig. 1 shows the result of an analysis of the humulone and lupulone groups of Hallertau hops. The concentration gradient of the eluant was accomplished by letting 90% (w/w) methanol flow into a constant volume mixing chamber loaded with 55% (w/w) methanol, the effluent of which was fed to the column. The resolution was satisfactory but somewhat less complete than with normal elution. Thus lupulone and adlupulone were not separated.

In other experiments it has been possible to vary the methanol concentration over still wider ranges. Carbon tetrachloride extracts of hops have been shown to be very heterogeneous mixtures containing substances which may be eluted at methanol percentages varying from 0 to 85% or still higher.

The collection of eluate fractions from a chromatographic column and the determination of the ultraviolet absorption of each fraction are inherently subject to systematic errors, which may be of a considerable magnitude, especially when working with substances which are unstable to air. The column effluent was therefore passed through a 5 mm absorption cell in a Hilger Uvispek spectrophotometer and the optical density at 280 or 325 mμ recorded automatically by a Brown recorder.


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On the Production of Magnetic Centres in Glycine by X-Rays

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The demonstration of the production of magnetic centres in a variety of chemicals by ionizing radiation might well lead to a better understanding of the mechanisms by which a relatively small amount of energy absorbed in the form of radiation causes marked biological effects. The paramagnetic resonance spectra obtained have been tentatively explained by assuming that organic free radicals constitute the magnetic centres. These results are of great interest to radiation biology when considered in relation to other evidence concerning aftereffects of irradiation and protection against radiation damage. We thought it worth while to try and obtain data in order to evaluate, more quantitatively, the importance of the earlier observations for radiation biology. As a first step we investigated the relation between the dose of radiation administered and the number of magnetic centres produced in an amino acid, glycine.

For irradiation we used X-rays produced at 175 kV and filtered by 2 mm of aluminium (about 800 r-min⁻¹). The dose was measured during the irradiation by means of a wavelength independent thimble chamber connected to a valve electrometer and integrating counter (Duplex-Dosemeter) carefully calibrated in Roentgens. The glycine, in amounts of about 150 mg, in the form of small crystals, was irradiated in evacuated and sealed quartz tubes, which were only half-filled. This en-

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Derivative of paramagnetic resonance spectrum of glycine, as recorded immediately after irradiation. Paramagnetic resonance spectrum of glycine, _immediately and_ 12 days after irradiation (integrated derivative curves).

Fig. 1 a. Derivative of paramagnetic resonance spectrum of glycine, as recorded immediately after irradiation. Paramagnetic resonance spectrum of glycine, immediately and _12 days after irradiation_ (integrated derivative curves).

abled one half of the tubes to be shielded from irradiation, and, after shaking the glycine into the unirradiated part of the tube, the measurement of paramagnetic resonance was conducted excluding completely the well known effects of radiation on quartz. The unirradiated quartz did not show any detectable signal. The spectrometer used contained a simple microwave bridge operating at a frequency of 9 400 Mc. The signal reflected from the cavity could be recorded as an absorption curve on an oscillograph or as a derivative curve by means of a pen recorder. Irradiation and magnetic measurements were conducted at room temperature.

The spectra observed immediately after the irradiation (Fig. 1 a) showed a central peak at a g-value of 2.004 (± 0.001) and two satellites at ± 22.4 gaus. On keeping the material in vacuo and repeating the measurements the shape of the spectrum was found to change slowly (Fig. 1 b), though the area under the absorption curves remained essentially constant. This may be taken to indicate a constant number of magnetic centres. The decay, if any, is less than 1 % per day at about 20° C. Spectra of identical shape could be recorded in the range from 2 kr to 350 kr, thus including much of the region of biological interest. Since the shape of the curves is independent of dosage, the linear rise in amplitude with dosage, shown in Fig. 2, indicates a corresponding increase in the number of magnetic centres.

Intensity, shape, and time dependence of the recorded spectra were not changed by the presence of air, neither during nor after irradiation, contrasting with results published for some other substances.

By comparing the areas under the absorption spectra of irradiated glycine and of a known amount of free radicals (diphenylpicryl hydrazyl) a rough estimate of the concentration of free radicals produced per unit dose in glycine could be obtained. The yield was found to be of the order of magnitude of 1 radical per 10 eV of radiation energy absorbed, i.e., much higher than the values reported for plastics.

Another series of experiments was conducted with living material. Embryos cut from resting barley seeds yielded a resonance peak at a g-value characteristic of free radicals, when irradiated in air or in nitrogen. A more detailed account of these studies will appear elsewhere.

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Fig. 2. Derivative amplitude (arbitrary units) of the paramagnetic absorption curve of glycine as a function of dose of irradiation by X-rays. X and O: two different series of experiments.

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The Reduction of Plutonium by Tetravalent Uranium

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In the search for substances to reduce small amounts of plutonium of higher oxidation states to Pu(III) in the presence of large amounts of U(VI), U(IV) has been tested under a number of various conditions. The use of U(IV) for reducing plutonium does not seem to have been described previously.

Tetravalent uranium at concentrations \( \gg 0.01 \text{ M} \) has been found to be stable in solutions of most acids for weeks or months if air is excluded. However, the stability in HNO₃ is poor, but increases with the U(IV) concentration. Maximum stability in HNO₃ seems to be obtained around 1 M HNO₃, where about 50 % of originally 0.1 M U(IV) is reduced in a little more than an hour at room temperature. Thus U(IV) can be used as a reducing agent even in HNO₃ solutions, provided the time of the experiments does not much exceed an hour.

In 1 M HNO₃, Pu(IV) and Pu(VI) are quantitatively (\( \approx 99 \%) \) reduced to Pu(III) by 0.1 M U(IV) within 5 min at room temperature. This effect can theoretically be used for separating uranium from plutonium in liquid-liquid extractions, if U(VI), U(IV), Pu(VI) and Pu(IV) prefer the organic phase and Pu(III) only prefers the aqueous phase. The practical usefulness of this was tested in two sets of experiments.

In the first set of experiments two solutions were prepared. The organic solution consisted of methyl isobutyl ketone, which was 1.0 M in HNO₃ and contained Pu(IV) and Pu(VI); the aqueous solution was 0.7 M in HNO₃, 3 M in Ca(NO₃)₂ and contained a reductant. When these two phases are equilibrated with each other, the distribution ratio of the metals between the organic and the aqueous phases is for U(VI) 12, U(IV) ~ 10, Pu(VI) 13, Pu(IV) 9, and Pu(III) \( \approx 0.001 \); of these species thus only Pu(III) prefers the aqueous phase, and can easily be separated from the other metal ions. After the two phases had been shaken together, the amount of reduced plutonium (i.e., Pu(III)) was determined. Some of the results obtained with different reductants are given in the table for 5 min equilibration time.

<table>
<thead>
<tr>
<th>Reducing agent</th>
<th>Pu reduced</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05 M NH₄OH · HCl + 0.005 M Fe³⁺</td>
<td>65 ± 8</td>
<td></td>
</tr>
<tr>
<td>0.05 M H₂NOSO₂NH₄ + 0.005 M Fe³⁺</td>
<td>65 ± 10</td>
<td></td>
</tr>
<tr>
<td>0.05 M U⁴⁺ in air</td>
<td>70 ± 10</td>
<td></td>
</tr>
<tr>
<td>0.05 M U⁴⁺ in N₂</td>
<td>85 ± 3</td>
<td></td>
</tr>
</tbody>
</table>

It is seen that U(IV) is equally or more effective than iron(II) plus hydroxylamine or sulfamiate.

In the second set of experiments, the organic phase contained Pu(IV) as the uncharged TTA complex in 0.1 M TTA (thienyltrifluoroacetone) in benzene, while the aqueous phase was 0.33 M in HClO₄ and contained U(IV). For 0.010 M U(IV), 14 % Pu had been reduced in 30 min, and 60 % in 20 h.

These experiments show the usefulness of U(IV) as a reducing agent for plutonium, either in aqueous solution or in two-phase systems.