A POSSIBLE PRIMARY QUANTUM CONVERSION ACT OF PHOTOSYNTHESIS

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To account for the observation that illumination prevents the appearance of newly assimilated carbon in the compounds of the tricarboxylic acid cycle, it was suggested that the light shifts the steady-state condition of the thioctic acid-containing coenzyme toward the reduced (dithiol) form, in which condition it is incapable of oxidatively decarboxylating pyruvic acid newly formed from CO₂ to give rise to the acetyl-CoA required to bring this carbon into the compounds of the Krebs cycle. We are here reporting some observations leading to the further suggestion that this shift toward the dithiol form is the direct result of the light action and that a biradical formed by dissociation of the disulfide bond in a strained five-membered disulfide containing ring (as in 6,8-thioctic acid and trimethylenedisulfide) is the species in which the quantum absorbed by the plant pigments and stored as electronic excitation in chlorophyll appears first as chemical bond potential energy; i.e., that a possible primary quantum conversion act of photosynthesis is represented by the equation

\[ \text{Chl}^* + S - S \rightarrow \text{Chl (ground)} + S + S \] (I)
Subsequent abstraction of H atoms$^{14,15}$ from a suitable donor by the thyl free radicals would lead to the dithiol which would be reoxidized ultimately by CO$_2$. The residual oxidation product of the H donor would lead in the end to molecular oxygen. It is obvious that on both the reductant and oxidant sides the chemical products of the conversion of several quanta will be required to accomplish the reduction of each CO$_2$ molecule and the generation of each O$_2$ molecule.

These subsequent reactions, being strictly chemical, may lead to diverse energy rearrangements. For example, the chemical potential of reduced carbon might be converted by oxidative phosphorylation reactions into the energy of phosphoric anhydrides, which in turn, could raise the potential energy of intermediates in the reaction sequences leading to the evolution of molecular oxygen and to the reduction of CO$_2$.$^{16,17}$

A value of the dissociation energy for this particular disulfide link lying in the region of 30-40 Kcal. would constitute not only permissive evidence for reaction (I) but positive evidence in its support, since hitherto it has been difficult to suggest any likely primary chemical step capable of usefully absorbing the greater part of the 30-40 Kcal. quantum of electronic excitation available for photosynthesis. Estimates of $D(\text{RS-\text{SR}})$ from simple open chain compounds range from $50_{18}^{12} \text{ to } 70_{10}^{10}$ Kcal. However, the fact that 5,8-thioctic acid is colorless, while 6,8-thioctic acid is yellow,$^{20}$ suggested that the incorporation of the S-S bond into a 5-membered ring might indeed introduce sufficient strain into it so as to reduce the S-S dissociation energy by as much as 25-30 Kcal., thus bringing it down into the required range. A number of experiments have been performed using the product of the reaction of Na$_2$S$_2$ with (CH$_2$)$_3$Br$_2$ (trimethylenedisulfide) as a model substance. Its absorption
spectrum together with that of the two thiocic acids and n-propyl disulfide is shown in Figure 1. A light induced fading of diphenylpicrylhydrazyl at room temperature and dependent upon the presence of the disulfide was demonstrated. A photochemical polymerization of the disulfide by light of wave length greater than 4,000 Å was sensitized by Zn tetraphenylporphin. These results may be taken to indicate that the dissociation energy of the disulfide bond in this compound is something less than 50 Kcals. and that this dissociation may be brought about by energy transfer from some other molecule in a suitably excited state.

That such energy transfers may take place, and especially efficiently in condensed systems, has been amply demonstrated. Since the grana have the optical properties of a condensed chlorophyll phase, a quantum absorbed anywhere within that phase is very rapidly transferred among the identical molecules of that phase at the singlet or the triplet levels. Concomitant with this, there occurs a decrease in the probability of emission as fluorescence and hence an increase in the availability of the quantum for chemical transformation (disulfide fission). In such a system, the high efficiency of energy conversion may be retained even though the ratio of chlorophyll to disulfide molecules be large ($10^2$-$10^3$).
References

(1) The work described in this paper was sponsored by the U. S. Atomic Energy Commission.

(2) Rockefeller Fellow, 1952-1953, while on leave from Brasenose College and the Dyson Perrins Laboratory, Oxford University, England.

(3) A. A. Benson and M. Calvin, J. Exptl. Bot., 1, 63 (1950).


(5) M. Calvin and Peter Massini, Experientia, in press.


(16) Phosphorus Metabolism, I, Johns Hopkins Press, 1951, Sec. V.


(20) Several milligrams of each of these synthetic$^{21}$ products were obtained through the courtesy of Dr. T. H. Jukes of Lederle Laboratories.


(32) D. McClure, private communication.

(33) We are indebted to Mr. Paul Hayes for the determination of these absorption spectra.
Caption to Figure

Figure 1 - Absorption spectra of several disulfides

[Graph showing absorption spectra of various disulfides in alcohol]