Orthomolecular Psychiatry -- Linus Pauling, Ph.D.

Varying the concentrations of substances normally present in the human body may control mental disease.

The methods principally used now for treating patients with mental disease are psychotherapy (psychoanalysis and related efforts to provide insight and to decrease environmental stress), chemotherapy (mainly with the use of powerful synthetic drugs, such as chlorpromazine, or powerful natural products from plants, such as reserpine), and convulsive or shock therapy (electroconvulsive therapy, insulin coma therapy, pentylenetetrazol shock therapy). I have reached the conclusion, through arguments summarized in the following paragraphs that another general method of treatment, which may be called orthomolecular therapy, may be found to be of great value, and may turn out to be the best method of treatment for many patients.

Orthomolecular psychiatric therapy is the treatment of mental disease by the provision of the optimum molecular environment for the mind, especially the optimum concentrations of substances normally present in the human body. An example is the treatment of phenylketonuric children by use of a diet containing a smaller than normal amount of the amino acid phenylalanine. Phenylketonuria results from a genetic defect that leads to a decreased amount or effectiveness of the enzyme catalyzing the oxidation of phenylalanine to tyrosine. The patients on a normal diet have in their tissues abnormally high concentrations of phenylalanine and some of its reaction products, which, possibly in conjunction with the decreased concentration of tyrosine, cause the mental and physical manifestations of the disease (mental deficiency, severe eczema, and others). A decrease in the amount of phenylalanine ingested results in an approximation to the normal or optimum concentrations and to the alleviation of the manifestations of the disease, both mental and physical.
The functioning of the brain is dependent on its composition and structure; that is, on the molecular environment of the mind. The presence in the brain of molecules of N, N-diethyl-D-lysergamide, mescaline, or some other schizophrenogenic substance is associated with profound psychic effects. Cherkin has recently pointed out that in 1799 Humphry Davy described similar subjective reactions to the inhalation of nitrous oxide. The phenomenon of general anesthesia also illustrates the dependence of the mind (consciousness, ephemeral memory) on its molecular environment.

The proper functioning of the mind is known to require the presence in the brain of molecules of many different substances. For example, mental disease usually associated with physical disease, results from a low concentration in the brain of any one of the following vitamins: thiamine (B1), nicotinic acid or nicotinamide (B3), pyridoxine (B6), cyanocobalamin (B12), biotin (H), ascorbic acid (C), and folic acid. There is evidence that mental function and behavior are also affected by changes in the concentration in the brain of any of a number of other substances that are normally present, such as L (+)-glutamic acid, uric acid, and gamma-aminobutyric acid.

**Optimum Molecular Concentrations**

Several arguments may be advanced in support of the thesis that the optimum molecular concentrations of substances normally present in the body may be different from the concentrations provided by the diet and the gene controlled synthetic mechanisms, and, for essential nutrilites (vitamins, essential amino acids, essential fatty acids) different from the minimum daily amounts required for life or the “recommended” (average) daily amounts suggested for good health. Some of these arguments are presented in the following paragraphs.

**Evolution and Natural Selection**

The process of evolution does not necessarily result in the normal provision of optimum molecular concentrations. Let us take ascorbic
acid as an example. Of the mammals that have been studied in this respect, the only species that have lost the power to synthesize ascorbic acid and that accordingly require it in the diet are man, other Primates (rhesus monkey, Formosan long-tail monkey, and ring-tail or brown capuchin monkey), the guinea pig, and an Indian fruit-eating bat (Pteropus medius).

Presumably the loss of the gene or genes controlling the synthesis of the enzyme or enzymes involved in the conversion of glucose to ascorbic acid occurred some 20 million years ago in the common ancestor of man and other Primates, and occurred independently for guinea pig and for one species of bat and one bird, in each case in an environment such that ascorbic acid was provided by the food. For a mutation rate of 1/20,000 per gene generation and for even a very small advantage for the mutant (0.01 percent more progeny) the mutant would replace the earlier genotype within about 1 million years. The advantage to the mutant of being rid of the ascorbic-acid-synthesis machinery (decrease in cell size and energy requirement, liberation of machinery for other purposes) might well be large, perhaps as much as 1 percent; a disadvantage nearly as large (less by 0.01 percent) resulting from a less than optimum supply of dietary ascorbic acid would not prevent the replacement of the earlier species by the mutant. Hence, even if the amount of the vitamin provided by the diet available at the time of the mutation were less than the optimum amount, the mutant might still be able to replace its predecessor. Moreover, it is possible that the environment has changed during the last 20 million years in such a way as to provide a decreased amount of the vitamin. Even a serious disadvantage of the changed environment would not lead to a mutation restoring the synthetic mechanism within a period of a few million years, because of the small probability of such mutations, far smaller than of those resulting in loss of function.

Fig. 1. Diagrammatic representation of growth rate or other vital property of an organism as function of the concentration of vital substance in the organism, showing the concentration at which the
differential advantage of an increased amount of vital substance is just balanced by the differential disadvantage resulting from an increased amount of machinery for synthesis, and the concentration that gives optimum functioning without consideration of the burden of the machinery for synthesis.

**Fig. 2** (right). The observed rate of growth of a pyridoxine-requiring Neurospora mutant, as function of the concentration of pyridoxine in the medium.

Moreover, the process of natural selection may be expected later on to lead to the survival of a species or strain that synthesizes somewhat less than the optimum amount of an autotrophic vital substance rather than of the species or strain that synthesizes the optimum amount. To synthesize the optimum amount requires almost double the biological machinery as synthesizing half the optimum amount. As suggested in Fig. 1, the evolutionary disadvantage of synthesizing a less than optimum amount of the vital substance may be small, and may be outweighed by the advantage of requiring a smaller amount of biological machinery. Evidence from the study of microorganisms is discussed in the following paragraphs.

**Evidence from Microbiological Genetics**
Many mutant microorganisms are known to require, as a supplement to the medium in which they are grown, a substance that is synthesized by the corresponding wild-type organism (the normal strain). An example is the pyridoxine-requiring mutant of Neurospora sitophila reported by G. W. Beadle and E. L. Tatum in their first Neurospora paper, published in 1941. Several species of Neurospora that have been extensively studied are known to be able to grow satisfactorily on synthetic media containing inorganic salts, an inorganic source of nitrogen, such as ammonium nitrate, a suitable source of carbon, such as sucrose, and the vitamin biotin. All other substances required by the organism are synthesized by it. Beadle and Tatum found that exposure to x-radiation produces mutant strains such that one substance must be added to the minimum medium in order to permit the growth at a rate approximating that of the normal strain. Their pyridoxine-requiring mutant was found to grow on the standard medium at a rate only 9 percent of that of the normal strain.

When pyridoxine (vitamin B6) is added to the medium, the rate of growth of this strain at first increases nearly linearly with the concentration of the added pyridoxine, and then increases less rapidly, as shown in Fig. 2. The growth rate of the normal strain without added pyridoxine is equal to that of the mutant with about 10 micrograms of the growth substance per liter in the medium. At a concentration about four times this value (40 micrograms per liter) the growth rate of the mutant strain reaches a value 7 percent greater than that of the normal strain without added pyridoxine.

The point of maximum curvature of the curve in Fig. 2, at about 3.2 micrograms of pyridoxine per liter (indicated by a cross), may be reasonably considered to mark the division between the region of vitamin deficiency (to the left) and the region of normal vitamin supply (to the right), such as might permit the mutant to compete with the wild type, which has the growth rate represented by the filled circle in Fig. 2. The point marked by the cross might well correspond to an “adequate” or “recommended” amount of the vitamin, in that the growth rate of the mutant is only 12 percent less than that of the wild strain, and that the amount of the vitamin would...

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have to be increased threefold to make up this 12 percent.

As shown in Fig. 2, quadrupling the concentration of pyridoxine that gives the mutant a growth rate equal to that of the wild type causes a further increase in growth rate by nearly 10 percent. The growth rates of the mutant and the wild type at very large concentrations of the vitamin have not been measured, so far as I know, and the optimum concentration is not known. From the work of Beadle and Tatum the optimum concentration may be taken to be greater than 40 micrograms per liter; that is, more than ten times the “adequate” concentration for the mutant and more than four times the concentration equivalent to the synthesizing capability of the wild type. The growth rate of the mutant at the optimum concentration is more than 22 percent greater than that at the “adequate” concentration and more than 9 percent greater than that of the normal strain.

Similar results have been reported for other mutants of Neurospora. The values found by Tatum and Beadle for a p-aminobenzoic acid-requiring mutant of Neurospora crassa as a function of the concentration of p-aminobenzoic acid added to the standard medium are shown in Fig. 3. The growth-rate curve is similar in shape to that for the pyridoxine-requiring mutant. The value of the growth rate for the normal strain of Neurospora crassa with no added p-aminobenzoic acid is equal to that for the mutant at a concentration of added p-aminobenzoic acid of about 15 micrograms per liter. A value about 4 percent greater is found for the normal strain at 40 micrograms per liter and for the mutant strain at 80 micrograms per liter, as indicated in Fig. 3.

**Fig. 3** (left). The observed rate of growth of a p-aminobenzoic-acid-requiring Neurospora mutant (Tatum and Beadle, 1942), as function of concentration of the growth substance in the medium. **Fig. 4** (right). Observed rate of growth of a para-aminobenzoic acid-requiring Neurospora mutant as function of the logarithm of the concentration of p-aminobenzoic acid.
It is customary to plot values of the growth rate against the logarithm of the concentration of the growth substance, as shown in Fig. 4. The amount of increase accompanying a doubling in the concentration of the growth substance is a maximum at 1.25 to 2.5 micrograms per litre, and decreases thereafter to about half the value for each successive doubling.

From these two examples we see that there may be a significant increase in rate of growth of the normal strain through addition of some of the growth substance that it synthesizes to the medium in which it is grown; that is, that the amount of the growth substance that is synthesized by the normal strain is not the optimum amount, but is somewhat less- approximately 7 percent less in the case of pyridoxine (with the normal strain of Neurospora sitophila) and 4 percent less for p-aminobenzoic acid (with the normal strain of Neurospora crassa). Many other examples are known of microorganisms that grow more abundantly in a medium containing vitamins, amino acids, or other substances that they are able to synthesize than on a minimum medium.

Evidence supporting the above arguments has been presented recently by Zamenhof and Eichhorn in a paper entitled “Study of microbial evolution through loss of biosynthetic functions: Establishment of 'defective' mutants.” These authors carried out experiments involving competitive growth in a chemostat of an auxotrophic mutant (a mutant requiring a nutrilite) and a
prototrophic parent in a medium of constant composition containing the nutrilite. They found that the “defective” mutant has a selective advantage over the prototrophic parental strain under these conditions. For example, an indole-requiring mutant of Bacillus subtilis was found to show a strong selective advantage over the prototrophic back-mutant when the two were grown together in a medium containing tryptophan; the relative number of cells of the latter decreased 10 fold in 54 generations. They also found that greater advantage to the auxotroph accompanies a greater number of biosynthetic steps that have been dispensed with (earlier block in a series of reactions), with the final metabolite available. They point out that a mutant with a gene deletion would be at a distinct selective advantage over a point mutant, in that not only the synthesis of the metabolite, but also that of the structural gene, the messenger RNA, and perhaps the inactive enzyme itself would be dispensed with, and that accordingly the mutant with a deletion would replace the point mutant in competition. They mention evidence that some of the “defective” strains occurring in nature have lost one or more of their structural genes by deletions, rather than by point mutations.

Molecular Concentrations and Rate of Reaction

Most of the chemical reactions that take place in living organisms are catalyzed by enzymes. The mechanisms of enzyme-catalyzed reactions in general involve (i) the formation of a complex between the enzyme and a substrate molecule and (ii) the decomposition of this complex to form the enzyme and the products of the reaction. The rate determining step is usually the decomposition of the complex to form the products, or, more precisely, the transition through an intermediate state of the complex, characterized by activation energy less than for the uncatalyzed reaction, to a complex of the enzyme and the products of reaction, with a rapid dissociation. Under conditions such that the concentration of the complex corresponds to equilibrium with the enzyme and the substrate, the rate of the reaction is given by the following equation [the Michaelis-Menten equation]:

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In this equation \([S]\) is the concentration of the substrate, \(E\) is the total concentration of enzyme (present both as free enzyme and enzyme complex), \(K\) is the equilibrium constant for formation of the enzyme complex \(ES\), and \(k\) is the reaction-rate constant for decomposition of the complex to form the enzyme and reaction products. This equation corresponds to the case in which there are no enzyme inhibitors present.

Values of the reaction rate calculated from this equation for different values of \(K\) are shown in Fig. S. The curves are similar in shape to those of Figs. 2 and 3. At concentrations much smaller than \(K\) (\(-1\)) the reaction rate is proportional to the concentration of substrate. At larger concentrations, as the amount of enzyme complex becomes comparable to the amount of free enzyme, the reaction rate changes from the linear dependence. At substrate concentration equal to \(K\) (\(-1\)) the slope of the curve is one-quarter of the initial slope, and the value is one-half of the value corresponding to saturation of the enzyme by the substrate.

The similarity of the curves of Figs. 2 and 3 to appropriate curves in Fig. 5 suggests that the growth substance may be involved in an enzyme-catalyzed reaction in which it serves as the substrate. The normal strain of the organism manufactures an amount of the substrate such as to permit the reaction to take place at what may be considered a normal rate, 90 or 95 percent of the maximum rate, which corresponds to saturation of the enzyme. As described above, the gain in reaction rate associated with the manufacture of a larger amount of the substrate, with a corresponding advantage to the organism, might be balanced by the disadvantage to the organism.
associated with the upkeep of the larger amount of machinery required to manufacture the increased amount of substrate. An increase in rate of this reaction could also be achieved by an increase in the amount of the enzyme synthesized by the organism. Here, again, the advantage to the organism resulting from this increase may be overcome by the disadvantage associated with the increase in the amount of machinery required for the increased synthesis. During the process of evolution there has presumably been selection of genes determining the concentrations of the enzymes catalyzing successive reactions such as to achieve an approximation to the optimum reaction rate with the smallest amount of disadvantage to the organism.

The rate of an enzyme-catalyzed reaction is approximately proportional to the concentration of the reactant, until concentrations that largely saturate the enzyme are reached. The saturating concentration is larger for a defective enzyme with decreased combining power for the substrate than for the normal enzyme. For such a defective enzyme the catalyzed reaction could be made to take place at or near its normal rate by an increase in the substrate concentration, as indicated in Fig. 5. The short horizontal lines intersecting the curves indicate what may be called the “normal” reaction rate, 80 percent of the maximum. For $K = 2$ the “normal” rate is achieved at substrate concentration $[S] = 2$. At this substrate concentration the reaction rate is only 29 percent of the maximum and 35 percent of “normal” for a mutated enzyme with $K = 0.2$; it could be raised to the “normal” value by a tenfold increase in the substrate concentration, to $[S] = 20$. Similarly, the still greater disadvantage of low reaction rate for a mutated enzyme with $K$ only 0.01 could be overcome by a 200-fold increase in substrate concentration, to $[S] = 400$. This mechanism of action of gene mutation is only one of several that lead to disadvantageous manifestations that could be overcome by an increase, perhaps a great increase, in the concentration of a vital substance in the body. These considerations obviously suggest a rationale for megavitamin therapy.
Molecular Concentrations and Mental Disease

The functioning of the brain and nervous tissue is more sensitively dependent on the rate of chemical reactions than the functioning of other organs and tissues. I believe that mental disease is for the most part caused by abnormal reaction rates, as determined by genetic constitution and diet, and by abnormal molecular concentrations of essential substances.

The operation of chance in the selection for the child of half of the complement of genes of the father and mother leads to bad as well as to good genotypes, and the selection of foods (and drugs) in a world that is undergoing rapid scientific and technological change may often be far from the best. Significant improvement in the mental health of many persons might be achieved by the provision of the optimum molecular concentrations of substances normally present in the human body.

Among these substances, the essential nutrients may be the most worthy of extensive research and more thorough clinical trial than they have yet received. An important example of an essential nutrient that is required for mental health is vitamin B12, cyanocobalamin. A deficiency of this vitamin, whatever its cause (pernicious anemia; infestation with the fish tapeworm Diphyllobothrium, whose high requirement for the vitamin results in deprivation for the host; excessive bacterial flora, also with a high vitamin requirement, as may develop in intestinal blind loops), leads to mental illness, often even more pronounced than the physical consequences. The mental illness associated with pernicious anemia [a genetic defect leading to deficiency of the intrinsic factor (a mucoprotein) in the gastric juice and the consequent decreased transport of cyanocobalamin into the blood] often is observed for several years in patients with this disease before any of the physical manifestations of the disease appear. A pathologically low concentration of cyanocobalamin in the serum of the blood has been reported to occur for a much larger fraction of patients with mental illness than for the general population. Edwin, www.healthoracle.org
Holten, Norum, Schrumpf, and Skaug determined the amount of B12 in the serum of every patient over 30 years old admitted to a mental hospital in Norway during a period of 1 year. Of the 396 patients, 5.8 percent (23) had a pathologically low concentration, less than 101 picograms per millilitre, and the concentration in 9.6 percent (38) was subnormal (101 to 150 picograms per millilitre). The normal concentration is 150 to 1300 picograms per millilitre.

The incidence of pathologically low and subnormal levels of B12 in the serums of these patients, 15.4 percent, is far greater than that in the general population, about 0.5 percent (estimated from the reported frequency of pernicious anemia in the area, 9.3 per 100,000 persons per year). Other investigators have also reported a higher incidence of low B12 concentrations in the serums of mental patients than in the population as a whole, and have suggested that B12 deficiency, whatever its origin, may lead to mental illness.

Nicotinic acid (niacin), when its use was introduced, cured hundreds of thousands of pellagra patients of their psychoses, as well as of the physical manifestations of their disease. For this purpose only small doses are required; the recommended daily allowance (National Research Council) is 12 milligrams per day (for a 70-kilogram male). In 1939 Cleckley, Sydenstricker, and Geeslin reported the successful treatment of 19 patients and in 1941 Sydenstricker and Cleckley reported similarly successful treatment of 29 patients with severe psychiatric symptoms by use of moderately large doses of nicotinic acid (0.3 to 1.5 grams per day). None of these patients had physical symptoms of pellagra or any other avitaminosis. More recently many other investigators have reported on the use of nicotinic acid and nicotinamide for the treatment of mental disease. Outstanding among them are Hoffer and Osmond, who since 1952 have advocated and used nicotinic acid in large doses in addition to the conventional therapy, for the treatment of schizophrenia. The dosage recommended by Hoffer is 3 to 18 grams per day, as determined by the response of the patient, of either nicotinic acid or nicotinamide, together with 3 grams per day of ascorbic acid.
Nicotinic acid and nicotinamide are nontoxic [the lethal dose, 50 percent effective (LD50), is not known for humans, but probably it is over 200 grams; the LD50 for rats is 7.0 grams per kilogram for nicotinic acid, and 1.7 grams per kilogram for nicotinamide], and their side effects, even in continued massive doses, seem not to be commonly serious. Among the advantages of nicotinic acid, summarized by Osmond and Hoffer, are the following: it is safe, cheap, and easy to administer, and it is a well-known substance that can be taken for years on end, if necessary, with only small probability of incidence of unfavorable side effects.

Another vitamin that has been used to some extent in the treatment of mental disease is ascorbic acid, vitamin C. A sometimes-recommended daily intake of ascorbic acid is 75 milligrams for healthy adults. Some investigators have estimated that the optimum intake is much larger: perhaps 3 to 15 grams per day, according to Stone. Williams and Deason have emphasized the variability of individual members of a species of animals; they have reported their observation of a 20-fold range of required intake of ascorbic acid by guinea pigs, and have suggested that human beings, who are less homogeneous, have a larger range.

Mental symptoms (depression) accompany the physical symptoms of vitamin-C deficiency disease (scurvy). In 1957 Akerfeldt reported that the serum of schizophrenics had been found to have greater power of oxidizing N-dimethyl-p-phenylenediamine than that of other persons. Several investigators then reported that this difference is due to a smaller concentration of ascorbic acid in the serum of schizophrenics than of other persons. This difference has been attributed to the poor diet and increased tendency to chronic infectious disease of the patients, and has also been interpreted as showing an increased rate of metabolism of ascorbic acid by the patients. It is my opinion, from the study of the literature, that many schizophrenics have an increased metabolism of ascorbic acid, presumably genetic in origin, and that the ingestion of massive amounts of ascorbic acid has some value in treating mental disease.

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Other vitamins (thiamine, pyridoxine, folic acid) and other substances [zinc ion, magnesium ion, uric acid, tryptophan, L (+)-glutamic acid, and others] influence the functioning of the brain. I shall review work on L (+)-glutamic acid as a further example.

L (+)-Glutamic acid is an amino acid that is present at rather high concentration in brain and nerve tissue and plays an essential role in the functioning of these tissues. It is normally ingested (in protein) in amounts of 5 to 10 grams per day. It is not toxic; large doses may cause increased motor activity and nausea. In 1944 Price, Waelsch, and Putnam reported favorable results for glutamic acid therapy of convulsive disorders [benefit to one out of three or four patients with petit mal epilepsy]. Zimmerman and Ross then reported an increase in maze-running learning ability of white rats given extra amounts of glutamic acid. Zimmerman and many other investigators then studied the effects of glutamic acid on the intelligence and behavior of persons with different degrees and kinds of mental retardation. L (+)-Glutamic is apparently more effective than its sodium or potassium salts. The effective dosage is usually between 10 and 20 grams per day (given in three doses with meals), and is adjusted to the patient as the amount somewhat less than that required to cause hyperactivity; improvement in personality and increase in intelligence (by 5 to 20 I.Q. points) have been reported for many patients with mild or moderate mental deficiency by several investigators.

**Fig. 5** (left). Curves showing calculated reaction rate RIR of catalyzed reaction as function of the concentration of the substrate, for different values of the equilibrium constant K for formation of the enzyme-substrate complex. **Fig. 6** (right). Values of the concentration of a vital substance in the blood and in the cerebrospinal fluid for three different assumed sets of value of blood-brain barrier permeability and rate of destruction in the cerebrospinal fluid.
Localized Cerebral Deficiency Diseases

The observation that the psychosis associated with pernicious anemia may manifest itself in a patient for several years before the other manifestations of this disease become noticeable has a reasonable explanation: the functioning of the brain and nervous tissue is probably more sensitively dependent on molecular composition than is that of other organs and tissues. The observed high incidence of cyanocobalamin deficiency in patients admitted to a mental hospital, mentioned above, suggests that mental disease may rather often be the result of this deficiency, and further suggests that other deficiencies in vital substances may be wholly or partly responsible for many cases of mental illness.

The foregoing arguments suggest the possibility that under certain circumstances a deficiency disease may be localized in the human body in such a way that only some of the manifestations usually associated with the disease are present. Let us consider, for example, an enzyme or other vital substance that is normally metabolized by the catalytic action of an enzyme normally present in the tissues and organs of the body. In a person of unusual genotype there might be an especially great concentration of this enzyme in one body organ, with essentially the normal amount in other organs. Through the action of this enzyme in especially great concentration the steady-state concentration of the vital substance in that organ might be
decreased to a level much lower than that required for normal function. Under these circumstances there would be present a deficiency disease restricted to that organ.

An especially important case is that of the brain. We may, as a rough model of the human body, consider two reservoirs of fluid, the blood and lymph, with volume $VI$, and cerebrospinal fluid, the extra cellular fluid of the brain and spinal column, with volume $V2$. We assume that a vital substance is destroyed in each of these reservoirs at a characteristic rate, corresponding to the rate constants $k1$ and $k2$, that it diffuses across the blood-brain barrier at a rate determined by the product of the permeability and area of the barrier and the difference $c2 - c1$ of the concentrations in the two reservoirs, and that it is introduced from the gastrointestinal tract into the first reservoir at a constant rate. The steadystate concentrations are then in the ratio $c1/c2 = 1 + (K2V2/PA)$

where $PA$ is the product of permeability and the area of the blood-brain barrier. The steadystate corresponds to the following system:

From this equation it is seen, as shown also in Fig. 6, that for small values of $k2V2/PA$ the difference in steadystate concentrations in the cerebrospinal fluid and the blood is small, but that through either decrease in permeability of the barrier or increase in the metabolic rate constant $k2$ the steadystate concentration in the brain becomes much less than that in the blood.

This simple argument leads us to the possibility of a localized cerebral avitaminosis or other localized cerebral deficiency disease. There is the possibility that some human beings have a sort of cerebral scurvy, www.healthoracle.org
without any of the other manifestations, or a sort of cerebral pellagra, or cerebral pernicious anemia. It was pointed out by Zuckerkandl and Pauling that every vitamin, every essential amino acid, every other essential nutrient represents a molecular disease which our distant ancestors learned to control, when it began to afflict them, by selecting a therapeutic diet, and which has continued to be kept under control in this way. The localized deficiency diseases described above are also molecular diseases, compound molecular diseases, involving not only the original lesion, the loss of the ability to synthesize the vital substance, but also another lesion, one that causes a decreased rate of transfer across a membrane, such as the blood-brain barrier, to the affected organ, or an increased rate of destruction of the vital substance in the organ, or wine other perturbing reaction.

It has been suggested by Huxley, Mayr, Osmond, and Hoffer, partially on the basis of the observations of Böök and Slater on the incidence of schizophrenia in relatives of schizophrenics, that schizophrenia is caused by a dominant gene with incomplete penetrance. They suggested that the penetrance, about 25 percent, may in some cases be determined by other genes and in some cases by the environment. I suggest that the other genes may in most cases be those that regulate the metabolism of vital substances, such as ascorbic acid, nicotinic acid or nicotinamide, pyridoxine, cyanocobalamin, and other substances mentioned above. The reported success in treating schizophrenia and other mental illnesses by use of massive doses of these vitamins may be the result of successful treatment of a localized cerebral deficiency disease involving the vital substances, leading to a decreased penetrance of the gene for schizophrenia. There is a possibility that the so-called gene for schizophrenia is itself a gene affecting the metabolism of one or another of these vital substances, or even of several vital substances, causing a multiple cerebral deficiency. I suggest that the orthomolecular treatment of mental disease, to be successful, should involve the thorough study of and attention to the individual, such as is customary in psychotherapy but less customary
in conventional chemotherapy. In the course of time it should be possible to develop a method of diagnosis (measurement of concentrations of vital substances) that could be used as the basis for determining the optimum molecular concentrations of vital substances for the individual patient and for indicating the appropriate therapeutic measures to be taken. My co-workers and I are carrying on some experimental studies suggested by the foregoing considerations, and hope to be able before long to communicate some of our results.

Summary

The functioning of the brain is affected by the molecular concentrations of many substances that are normally present in the brain. The optimum concentrations of these substances for a person may differ greatly from the concentrations provided by his normal diet and genetic machinery. Biochemical and genetic arguments support the idea that orthomolecular therapy, the provision for the individual person of the optimum concentrations of important normal constituents of the brain, may be the preferred treatment for many mentally ill patients. Mental symptoms of avitaminosis sometimes are observed long before any physical symptoms appear. It is likely that the brain is more sensitive to changes in concentration of vital substances than are other organs and tissues. Moreover, there is the possibility that for some persons the cerebrospinal concentration of a vital substance may be grossly low at the same time that the concentration in the blood and lymph is essentially normal. A physiological abnormality such as decreased permeability of the blood-brain barrier for the vital substance or increased rate of metabolism of the substance in the brain may lead to a cerebral deficiency and to a mental disease. Diseases of this sort may be called localized cerebral deficiency diseases. It is suggested that the genes responsible for abnormalities (deficiencies) in the concentration of vital substances in the brain may be responsible for increased penetrance of the postulated gene for schizophrenia and that the so-
called gene for schizophrenia may itself -be a gene that leads to a localized cerebral deficiency in one or more vital substances.