Immunoglobulin loci. Four H-chain gene loci, Ig-1, Ig-2, Ig-3 and Ig-4, have been found in the mouse, corresponding to the H-chains in the classes \( \gamma_{G_2a} \), \( \gamma_A \), \( \gamma_{G_2b} \) and \( \gamma_{G_1} \) respectively. The alleles at Ig-1 through Ig-3 were first defined serologically with alloantisera.

Detection of the reaction of alloantisera with immunoglobulin was by immunodiffusion in agar or, where more sensitive detection methods were necessary, by inhibition of precipitation of radioactive labeled antigens. The latter method was used for most of our analyses of allotypic differences.

Among some 70 inbred mouse strains, there are 8 alleles at the Ig-1 locus. These alleles display 11 distinct antigenic specificities on \( \gamma_{G_2a} \) immunoglobulins. Each allotype is cross reacting and is determined by various combinations of relatively few specificities.

Three other immunoglobulin loci closely linked to Ig-1 have been uncovered. Ig-2, Ig-3 and Ig-4 determine \( \gamma_A \), \( \gamma_{G_2b} \) and \( \gamma_{G_1} \) H-chains respectively. Ig-2 and Ig-3 were demonstrated by the same methods as Ig-1; i.e., cross reacting specificities found with alloantisera.

A fourth linked H-chain locus, Ig-4, has been defined, using allelic differences in electrophoretic mobility of \( \gamma_{G_1} \) immunoglobulins and Fc fragments. Thus, 4 of the 5 H-chains in mice are controlled by closely linked genes (forming the H-chain chromosome region).

Allelic electrophoretic mobility differences have been found for the previously identified H-chain loci, Ig-1 through Ig-3 as well. One or more of the serologically defined allotypes at each locus is strictly
associated with each electrophoretic mobility allotype. The use of electrophoretic mobility for allotype detection is new and of potentially great use for genetic studies of immunoglobulins.

Three heavy-chain allotypic specificities have been found on two of the mouse immunoglobulin classes, $\gamma G_{2a}$ and $\gamma G_{2b}$. The other described immunoglobulin allotypic specificities in the mouse are restricted to one or another H-chain class. This work helps understand the possibly evolutionary origins of shared or common specificities and is of importance to theories of generation of diversity in immunoglobulins.

Detection of Allotypic Antigens with Heterologous Antisera. Rabbits were immunized with either normal mouse immunoglobulins or isolated mouse myeloma proteins from BALB/c or (BALB/c X NZB)$F_1$ plasma cell tumors. The antisera were tested for antibodies directed to allotypic antigenic specificities by the method of inhibition of precipitation of $I^{125}$ labeled antigens.

Three $\gamma G_{2a}$ and one $\gamma G_{2b}$ allotypic specificities were detected with one or another of these rabbit antisera. These specificities each corresponded in mouse strain distribution with one of the allotypic specificities previously defined through the use of mouse isoantisera.

Plasma Cell Tumors. A new series of 15 myeloma proteins from plasma-cytomas induced in the Walter & Eliza Hall Institute, Melbourne, Australia, has been characterized here as to immunoglobulin type, physicochemical properties and allotypic specificities. Several of these tumors are now
L. A. Herzenberg - Immunogenetics and somatic cell genetics. (continued)

being maintained in this laboratory and are under further study. These are
the first plasmacytomas available from (BALB/c X NZB)F₁ hybrid mice hetero-
yzygous for allotype. Since they each produce large quantities of relatively homogeneous immunoglobulin, they greatly facilitate analysis of the
genetic control of immunoglobulins.

Three γG₂ myeloma proteins of plasma cell tumors induced in the
(NZB X BALB/c)F₁ mice have been analyzed for the isoantigens they carry. NZB mice are genotypically Ig-le Ig-3e, while BALB/c are Ig-la Ig-3a. Two
of the myeloma proteins are γG₂a globulins. One of these, GPC-7, carries
all the isoantigenic specificities of the Ig-1e allele while the other, GPC-8, carries all the isoantigenic specificities of the Ig-1a allele.
Thus, only one of the parental alleles of the mouse in which the tumor arose is expressed in each of these myeloma proteins.

The third myeloma protein, GPC-5, also carries the antigens of only
one parental strain (NZB). However, GPC-5, a γG₂b globulin, carries only
one of the Ig-3 specificities normally associated with γG₂b globulins of NZB. Most remarkably, it also carries one Ig-1 specificity normally
associated with γG₂a globulins of NZB. This is the first analysed mouse
myeloma shown (a) to express some but not all the antigenic specificities
normally associated with an allele, and (b) to carry antigenic specifici-
cities controlled by two distinct immunoglobulin loci.

Transplantation studies using immunoglobulin marked congenic strains.
Over the past three years, we have been breeding two pairs of strains of
mice such that the members of each pair would approach being congenic
(having the same genotype) except for the Ig-chromosome region. Skin
grafting and lymphoid cell transfer studies have now been carried out
with the first pair. The two strains involved are an inbred strain,
C3H.SW (CSW), and its congenic partner, CWR/5, which was derived by 5
backcrosses to CSW and 6 brother X sister matings. CSW has the Ig-1<sup>a</sup>
allele, while CWR has the Ig-1<sup>b</sup> allele. These strains are now completely
histocompatible for skin. Skin grafts, even after prior sensitization,
are permanently accepted in both directions. The production of donor
type immunoglobulins by transferred lymphoid cells was obtained with much
greater ease than in our previous experiments with allogenic strain cell
transfers. However, two barriers to the transferred cells were seen:
limited biological space and, what is more likely, a residual histo-
compatibility antigen(s) for lymphoid cells (although not for skin).
These barriers were overcome by limited sublethal X-irradiation and/or by
increased numbers of transferred cells. A most exciting prospect is the
finding of readily detectable amounts of donor globulins several weeks
after transfer of as few as 1,000 spleen cells.

Spleen, bone marrow and fetal liver, but not thymus cells, produce
γ-globulins for many weeks after injection into 600R irradiated congenic
recipients. As donor type levels increased, host type decreased in some
animals to very low levels.

These congenic strains are being used for studies of the ontogeny and
regulation of immunoglobulin production and for studies of the genetics of
antibody induction by specific antigens.

The congenic strains have found an important use in showing that fetal liver cells transferred into intact or thymectomized, sublethally irradiated hosts differentiate to produce donor allotype immunoglobulins. However, the thymectomized chimeras are unable to make specific antibodies to at least some antigens while the non-thymectomized controls do make antibodies when challenged.

Backcrossing of CWB/5 to CSW has been continued and a line is now being obtained by inbreeding at the 8th backcross to see whether the histoincompatibility has been bred out. Eleven backcrosses in all have been completed.

The other pair of congenic strains is C57BL/10 (Ig-1b) and its congenic partner, BTA (Ig-1a). A line is being developed by inbreeding after 11 backcross generations. It has not yet been tested for cell transfer.

Suppression of allotype production in young animals. Our recent work on the problem of allotype suppression is of importance in understanding the regulation of immunoglobulin production. In the mouse, an antibody directed against an immunoglobulin allotype, Ig-1b, passed from mother to offspring or injected into neonates, suppresses synthesis of immunoglobulin carrying Ig-1b. In allotype homozygotes as well as heterozygotes, the allotype suppression is manifested both by a delay of several weeks in attaining initial detectable allotype levels and a reduction in allotype level continuing into adulthood. There is evidence for a strong intralitter (as opposed to inter-litter) correlation of age of onset of immunoglobulin allotype synthesis.
H-2 antigen of NZB. Mice of the NZB strain develop an "auto-immune" disease with many similarities to the human disease, systemic lupus erythematosus. For transplantation studies, it is useful to know what H-2 type it is. We have shown that the NZB strain has the allele H-2\textsuperscript{d} both by hemagglutination and skin grafting studies. Snell's F\textsubscript{1} and component tests were used.
Leonard A. Herzenberg, Ph.D.
Associate Professor

Senior Research Associate

Leonore A. Herzenberg - suppression of immunoglobulin production, allotypic specificities, antiserum definition.

Postdoctoral Fellows

Ethel Jacobson - cell transfer studies with congenic mice.
Alma Luzzati (Research Associate) - single cell allotype studies.
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Predoctoral Fellows

Timothy Coburn - cell separation studies.
Peter Dolinger - antibody avidity studies.
Michel Facon - purification and structure studies on immunoglobulins.
Dale Hattis - isopyknic centrifugation of immunoglobulins.
Roy Riblet - lysozyme-producing mouse tumor and lysozyme polymorphism.
William Clewell (Medical Student) - isopyknic centrifugation of immunoglobulins.

Research Assistants

Priscilla Gibbs - primarily cell transfer and suppression of immunoglobulin synthesis.
Derek Hewgill - primary assay development and antiserum production.
CURRICULUM VITAE – LEONARD A. HERZENBERG

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Education
  1948 - 52 A. B. Brooklyn College, New York
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Publications
  See the attached sheets.
Molecular neurobiology. This project consists of basic research into the biochemistry of nerve cells particularly those aspects which differentiate nerve from other types of cells. Current research is therefore concerned with the properties and functions of macromolecules (a) which are specific to nerve cells, (b) whose mutant forms are associated with neurological disease and (c) which although found in other tissues interact specifically with nerve cells. An example of the latter is the Nerve Growth Factor which is an excellent model system for studying growth and differentiation in the nervous system.

The Nerve Growth Factor Protein. The Nerve Growth Factor (NGF) discovered by Levi-Montalcini is a protein which specifically stimulates the growth of functional axons from neurons of the sympathetic and embryonic sensory ganglia. Earlier work showed that the interaction of this protein with the target cell resulted in the stimulation of many biosynthetic pathways of which the stimulation of RNA synthesis appeared to be first in temporal sequence. Because fluorescent antibody to NGF interacted with components within the receptive neuron, it was concluded that part or all of the NGF molecule entered the cell. Nothing else is known about the details of the NGF interaction with the responsive neuron and our approach has been first to characterize the NGF protein itself in greater detail before going on to the metabolic experiments. It turns out that the NGF protein is an unusual protein. It is found in snake venoms and the adult male mouse salivary gland. A new NGF form has been purified forty fold from homogenates of the mouse gland by a procedure involving only gel filtration on Sephadex G-100, DEAE-chromatography and a second gel
filtration on Sephadex G-150. In the final product the NGF activity is associated, as judged by a number of physical criteria, with a single protein component comprising over 95% of the total protein of the fraction. It represents 2% of the soluble protein of the gland and 80% of the expressed activity of the gland homogenate. The molecular weight of this new NGF species is approximately 140,000 compared to 20-40,000 for NGF made by the older procedures. The large molecule is stable only between pH 5 and 8. Outside this pH range, it dissociates reversibly into smaller species with no, or considerably lower activity. The products of the complete dissociation are three groups of subunits, one acidic (α), one basic (β) and the third intermediate in net charge (γ), all three having molecular weights of approximately 30,000. The three groups of subunits have been isolated by chromatography on CM-cellulose at low pH and each displays heterogeneity on electrophoresis. Only the β subunit elicits a nerve growth factor type of response in the standard bioassay. It accounts for 25% or less of the original activity and is markedly unstable. However, the original highly active nerve growth factor protein is spontaneously reconstituted when the three subunits are mixed at neutral pH. All three types of subunits are needed for this regeneration; α and γ subunits, while not reacting with each other, combine separately with the β subunits but the resulting complexes have the same low activity and instability as the isolated β subunit itself.

Of the three types of subunits produced by the acid or alkaline dissociation of the 7S species of the mouse nerve growth factor protein, two of them
display heterogeneity on electrophoresis. The α subunits contain three major and one minor component and the γ subunits three components. Since each subunit component has a molecular weight around 30,000, they are not all derived from a single 7S nerve growth factor protein. Both individual α and γ subunits are separable by ion exchange chromatography and remain stable after separation. Recombination of any one α and any one γ subunit with the biologically active β subunit produces a 7S species with the physicochemical properties and increased biological activity characteristic of the original preparation of the nerve growth factor protein. The 7S species produced from a common γ but differing α subunits show small differences in electrophoretic mobility which reflect the mobility differences between the α subunits. When a common α but differing γ subunits are used in the recombination the resultant 7S species have the same mobility. Dissociation at either acid or alkaline pH of the 7S species formed from individual α and γ subunits produces only those subunits used in the initial recombination. These results suggest that the nerve growth factor protein preparation contains multiple forms of the 7S species all with the same general subunit composition but differing in the types of subunit they contain. In agreement with this hypothesis, the α subunit composition of the nerve growth factor protein is not constant across its migrating zone on electrophoresis or during elution from ion exchange resin but shows a continuous change from species containing predominantly the α subunits of higher mobility. The finding that the γ subunit composition also varies in the same way suggests that the multiple
forms of the nerve growth factor protein may be in equilibrium.

The NGF protein and its subunits have been successfully iodinated with $^{125}$I at low levels. These materials will be used to study both the subunit equilibria and, by autoradiographic techniques, the interaction of NGF with the receptive neurons. The nature of the differences between individual subunits of one class, the numbers of polypeptide chains within a subunit and the amino acid sequences and the identity of the non-protein moiety of NGF are all currently under investigation.

The high molecular weight 7S form of NGF exhibits another activity besides its growth promoting properties. It has a similar level of esterase activity on BAEE as does trypsin but a much lower level of proteolytic activity. These enzymatic activities are a property of the $\gamma$ subunits, the $\beta$ subunits, which are the only 7S NGF subunits to elicit a response in the bioassay, and the $\alpha$ subunits being inactive. The specific activities of the $\gamma^1$, $\gamma^2$ and $\gamma^3$ subunits are identical and higher than that of 7S NGF. The hydrolysis of BAEE by the $\gamma$ subunits proceeds in a linear manner in contrast to 7S NGF which shows an initial lag phase. Thus both the biological activity of the $\beta$ subunits and the enzymatic activity of the $\gamma$ subunits are altered by their interactions with the other two NGF subunits. Future work will by attempting to find specific inhibition of the $\gamma$ enzyme activity examine the relationship between enzymatic and biological activity.

Chemical Ontogeny and Polymorphisms of Nervous System Proteins. The major fraction of the protein of the nervous system is imbedded in a matrix
with lipid and carbohydrate and because of this inaccessibility has been relatively little studied. An investigation has therefore been made of the conditions required for the solubilization of water insoluble nervous tissue proteins and for their subsequent fractionation. The complete scheme for handling both water soluble and insoluble proteins may be briefly outlined.

Homogenization of whole brain in isotonic sucrose solution releases 15% of the total protein in water-soluble form. Subsequent exposure of the insoluble subcellular pellet to hypotonic buffer solution and ultrasonic treatment liberates an additional 20% of the total protein. Although several major differences were observed, these two aqueous extracts are similar in protein composition, being resolved into 50 distinct bands. Efficient dispersion of the insoluble residue remaining after osmotic shock, in buffer containing the non-ionic detergent Triton X-100 solubilizes an additional 34% of the total protein. By electrophoresis in the presence of Triton, this extract was fractionated into 10-15 protein bands, 3 or 4 of which were specific to this fraction and were profoundly influenced by the ratio of protein to detergent in the system. Electrophoresis of such extracts in the presence of sodium lauryl sulfate improves the resolution greatly, giving 20 clearly defined protein bands. Extraction of the Triton X-100-insoluble residue with the anionic detergent sodium lauryl sulfate results in solubilization of 21-31% of the total brain protein. By electrophoresis in the presence of sodium lauryl sulfate, this extract was fractionated into 40 distinct protein bands, the resolution of which are dependent upon the detergent concentration in
E. M. Shooter - Molecular neurobiology, genetic control of hemoglobin synthesis.
(continued)

the electrophoresis system. The four sequential protein extracts obtained under appropriate conditions with isotonic sucrose, hypotonic buffer, Triton X-100 and sodium lauryl sulfate represent protein pools of finite size and together account for at least 90% of the total brain protein. By electrophoretic analysis the majority of the proteins in the water soluble pools are similar but those in the Triton and sodium lauryl sulfate extracts differ not only from the water soluble proteins but from each other.

These procedures have been employed for investigation of the quantitative and qualitative changes in brain proteins during the ontogenetic development of the mouse. The most pronounced accumulation of proteins occurs during the initial 2 or 3 weeks after birth, with the greatest increases taking place among the detergent-soluble proteins, which account for a steadily increasing proportion of the total protein, indicating an increased synthesis of membranous and structural proteins. After the first few postnatal weeks, the changes in the protein content of brain are much less pronounced, occurring at a greatly diminished rate. During this time, the brain weight approaches a constant adult level, as do the quantities of protein solubilized by aqueous, Triton X-100, and 0.1% sodium lauryl sulfate solutions. However, the 1% sodium lauryl sulfate extract continues to increase markedly in protein content throughout the life span of the mouse, reaching a value of 20% of the total protein by 2 years after birth. These quantitative changes are accompanied by concurrent changes in the electrophoretic composition of each fraction, indicated by the appearance or disappearance of specific proteins or
shifts in their relative proportions, which occur at specific, although not identical, stages of development. The periods during which these changes in brain protein occurred correlate well with the known temporal sequence of brain histological and electrophysiological maturation and of behavioral development, implying a possible interrelationship of these phenomena.
E. M. Shooter - Molecular neurobiology, genetic control of hemoglobin synthesis
(continued)

Personnel

Senior (on sabbatical leave)

E. Glassman, Professor of Genetics
University of North Carolina - Molecular neurobiology (1968-69)

Sarane T. Bowen, Associate Professor of
Biology, San Francisco State - Genetic, developmental and
structural aspects of hemoglobins

Postdoctoral

M. Baker - Nerve growth factor - physical properties and relationship
of enzymatic and biological activities

H. R. Fisk - Nerve growth factor - molecular composition and equilibria,
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C. Louis - Physical and chemical properties of membrane proteins and
their interaction with lipid

T. Waehneldt - Micro methods for characterizing membrane proteins, sub-
cellular and developmental aspect with neurological mutants

Predoctoral

K. Borden - Specific acidic proteins in nerve cells

S. Morris - Turnover of brain proteins, axonal flow and synaptic
membrane production

R. Perez - Developmental aspects of brain proteins

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ences between subunits and sequence work

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1942-45 Natural Sciences Tripos (Part II in Chemistry)
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1942 Exhibitioner of Gonville & Caius College, Cambridge

1943 Minor Scholar of same.

1945 B.A. (Cantab.)

1945-46 and 1946-49 Research under Professor Sir Eric Rideal in the Department of Colloid Science, Cambridge and the Davy Faraday Laboratory of the Royal Institution, London (Proteins of the ground nut).

1949 M.A. (Cantab.)

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1948-50 Postdoctoral Fellowship with Dr. J. W. Williams, Department of Chemistry, University of Wisconsin, Madison, and partly with Dr. D. E. Green, Enzyme Institute, University of Wisconsin. (Enzymes of the electron transport system).

1950-53 Senior Scientist in charge of Biochemistry, Brewing Industry Research Foundation, Nutfield (Proteins and enzymes of barley and other brewing materials).

1953-63 Lecturer in Biochemistry, Department of Biochemistry, University College, London with Professor Ernest Baldwin. (Molecular biology of normal and abnormal haemoglobins; protein-ion interactions of ribonuclease).

1961-62 U.S.P.H.S. International Fellow, Department of Biochemistry, Stanford University School of Medicine, with Professor R. L. Baldwin (Replication of DNA).

1963-68 Associate Professor of Genetics, Stanford University School of Medicine (Molecular Neurobiology). Head of Neurobiology Group, Lt. Joseph P. Kennedy, Jr. Laboratories for Molecular Medicine.

1964 D.Sc. University of London (awarded for distinguished work in the field of Biochemistry).

1968-present Professor of Genetics, Stanford University School of Medicine.
J. Lederberg and E. C. Levinthal - Relevance of Current Program of Instrumentation Research Laboratory to Problems of Molecular Biology.

The activities of the Instrumentation Research Laboratory are largely supported by NASA in connection with their interest in an automated biological laboratory for the exploration of the planets. This has related to other biological interests in this department and elsewhere in the medical school in the areas of mass spectrometry, cell separation and classification, and the general question of computer managed instrumentation.

A. Mass Spectrometry

The connection to mass spectrometry is not too surprising since the same reasons that make mass spectrometry a powerful tool for biological explorations carried out remotely on a planet forty million miles distant from the earth make it a most sensitive and selective method for analyzing organic molecules important in problems of molecular biology germane to modern medicine. It is a potentially powerful method for investigating the unknown structure of unknown molecules in nerve tissue that form the engram which is part of the process of memory as well as Martian surface material. Cytochemistry via mass spectrometry is still a distant and challenging goal. However, significant progress has been made during this period. In addition to building a base for further advances our efforts have yielded results of present value.

The problem and the program can be subdivided into separate areas of concern. First, there is the question of volatilizing the molecules of interest. This can be approached by means of chemical modification of the class of molecules under investigation. We have successfully applied this concept to problems of resolution and identification of optical isomers of amino acids using the combination of a gas chromatograph and mass spectrometer. Molecules of biological interest are characterized by asymmetries at one or more of the carbon atoms incorporated in the molecule. From the viewpoint of the exobiologist this statement is the basis of the well-known significance of
optical activity as a clue for the recognition of life. The preparation of volatile diastereoisomers, their separation by gas chromatograph and their further identification by a mass spectrometer provides a method important to both terrestrial and extraterrestrial biology. The general concept is elucidated in the papers enumerated in the bibliography. Initially the technique was applied to the high sensitivity scanning of amino acids for optical activity while other work has demonstrated the general applicability of the method. A second but much more difficult method, which has the advantage that it is more directly applicable to the goal of cytochemistry, could conceivably utilize electron, heavy particle or photon beam energy. We have investigated the use of both heavy particles and laser photon beams. In the case of the former, we had useful but discouraging results. In the latter case our present results show some real promise.

A second subdivision of this effort addresses itself to acquiring basic data on the mass spectra of a large number of monomers of biological interest. A report covering the work on amino acids has already been published. Work on nucleotides and related products is in progress.

Computer control of mass spectrometers describes the third categorization of the program. Full advantage of a mass spectrometer as a biological tool can only be achieved when the instrument is under computer control. The typical processes of calibration and optimization of operating parameters are sufficiently complex that they require automation if it is desired to analyze a large number of spectra in a short period of time. We have designed and built a very effective system for computer operation of both a time-of-flight and quadrupole mass spectrometer. This system has been and is continuing to be used for biological research purposes. The system is being elaborated to
to provide a more sophisticated level of control. In addition, work is under-
way to achieve some degree of control of high resolution mass spectrometers
than use magnetic filters.

Fourthly, there is the question of data retrieval for subsequent com-
puter analysis. The ultimate goal of a two dimensional micro-description of
the distribution of molecules in a tissue by means of their mass spectra
presents formidable problems of data handling. The bandwidth requirements are
at least an order of magnitude greater than color video. High bandwidth data
retrieval and buffer storage are required. We have not directly confronted
this problem. General advances in the technology of high speed solid state
switching devices and information storage methods lend some hope for the
future. We have, however, made some modest steps. We have implemented an
interface system for a direct data link from a high resolution mass spectro-
meter to an IBM 360/50 computer.

The fifth and last subdivision of the program really represents most
clearly the ultimate goal for which the previously described efforts provide
the technological tools. Ultimately the spectra acquired must be analyzed.
This requires computer manipulation of chemical hypotheses. This poses a pro-
blem in both artificial intelligence and organic chemistry. A great deal of
progress has been made in this direction. Most of this research is supported
by the Advanced Research Projects Agency of the Office of the Secretary of
Defense, Contract No. SD-183, and carried out in collaboration with Professor
E. Feigenbaum of the Department of Computer Science.
While the high resolution mass spectrometer is perhaps the most capable single instrument for organic structural analysis, the sheer volume of its signal output poses formidable problems of data reduction and data analysis. These problems would be multiplied by the number of samples that would need to be processed by the micro-scanning mass spectrometer which is our ultimate goal. At one level, the problem is the identification of mass numbers with compositional formulas. However, no mass spectral signal is free of noise and great effort must then be spent to obtain an accurate determination of mass to ultimate resolution. Much of this effort is wasted when it does not answer a concrete question, i.e., which of a set of possible compositions is indicated by a given measurement. Even for all compositions, the corresponding mass numbers are not continuously distributed; they are rather the discrete set of numbers calculated from linear integral sums of nuclidic masses, and represented in the tables.

The tabulations and calculation programs (see bibliography) are the first step in a control program for the mass spectrometer. As soon as the peak is identified within a given mass neighborhood, the competing possibilities should be computed, then weighted in accordance with any other available information. This allows the experimental problem to be restated as a choice among competing possibilities, and the signal information need be accumulated only long enough to lead to a meaningful choice among them.

In solving a structure, the chemist hypothesizes a series of trial structures, then matches them with the data (in this case a mass spectrum, but this can be generalized to any data set) and accepts or rejects his trial
solutions, usually part by part, in a structure. Much of this tedious effort could be emulated or at least assisted by the computer in a program we call "mechanized induction".

For this purpose, a language has been devised for representing chemical structures in easily computable form; "Dendral '64". The development of this language required the filling of a surprising gap; the systematic application of simple topological principles to the field of chemical graphs - that is, a symbolic representation of organic molecules. Existing notations were found to be quite defective as the chemist already knows too well from his difficulties with nomenclature (other organizations like Chemical Abstract Service also recognize the problem and are working on it, but tend to compromise topological rigor for the benefit of established traditions in notation).

At any rate, with the help of some theorems on canonical forms of trees, and on Hamilton circuits of planar maps (for acyclic and cyclic structures respectively), a complete system has been worked out. This gives an algorithm by which the computer can generate an exact list of all isomers in a given composition.

By itself this is a futile approach to any but the simplest problems, since the number of possible isomers quickly exceeds the range of a fast computer. Heuristic and symbiotic methods are therefore called for whereby the computer emulates or cooperates in the use of human problem solving techniques in searching wisely selected parts of the space of possible solutions.

Professor Edward Feigenbaum and Dr. Richard Watson of the Computer Science Department participated in a cooperative effort to program efficient displays of structural ideas for conversational interaction with the computer. This
was a step to evaluate the chemists problem solving heuristics incorporating them in the machine program. The effort was limited at present by the existing computer facilities (a PDP-1 machine) with inadequate displays.

A complete description of the current status is given in an article by J. Lederberg and E. A. Feigenbaum. The following is the abstract of that paper.

"A computer program for formulating hypotheses in the area of organic chemistry is described from two standpoints: artificial intelligence and organic chemistry. The Dendral Algorithm for uniquely representing and ordering chemical structures defines the hypothesis-space; but heuristic search through the space is necessary because of its size. Both the algorithm and the heuristics are described explicitly but without reference to the LISP code in which these mechanisms are programmed. Within the program some use has been made of man-machine interaction, pattern recognition, learning, and tree-pruning heuristics as well as chemical heuristics which allow the program to focus its attention on a subproblem and to rank the hypotheses in order of plausibility. The current performance of the program is illustrated with selected examples of actual computer output showing both its algorithmic and heuristic aspects. In addition some of the more important planned modifications are discussed."
J. Lederberg and E. C. Levinthal - Relevance of Current Program of Instrumentation Research Laboratory to Problems of Molecular Biology.

B. Cell Separation

The work on cell separation has proceeded to the point where there are now three instruments capable of carrying out worthwhile biological experiments. The first of these is a volumetric cell separator, the second is a high speed fluorescent cell separator and the third is a cell separator and identifier developed by the Watson Labs of IBM under the direction of Dr. L. A. Kamentsky. While the ultimate goal of the Instrumentation Laboratory is the possible application of these principles to the biological exploration of the planets, our present efforts are directed toward current biological and medical problems. This has made it possible for us to gain the active participation of other scientists here at the Medical School. These have included Professor H. S. Kaplan, Executive Head of the Radiology and Radiotherapy; Dr. I. L. Weissman, Department of Radiology; Professor George Hahn, Department of Radiology; Professor Leonard Herzenberg, Department of Genetics; Professor R. Kallman, Department of Radiology.

Initially, we have established criteria for cell types which could be used as assay material while developing the prototype instruments leading to uses of instruments designed for cell separation in experiments that are biologically significant.

These criteria are:

1) Cells of at least two major size categories.
2) Cells which are not adherent to one another and which can be maintained in single cell suspension.