Drug discovery has been undergoing revolutionary changes for the last 15 years. The first change was in the shift from whole animal testing of compounds, to mechanistic based in vitro screening. This was based on new knowledge about molecular mechanisms brought about by advances in biochemistry. Some companies began to adapt in vitro mechanistic assays for “high throughput” screening, on the order of 10,000 to 20,000 compounds per year. At the same time, medicinal chemists began to recognize the advances being made in the understanding of the structure of enzymes and proteins, as well as to use x-ray to define the structure of the small molecules they had synthesized which exhibited desired biological activity. This approach enabled chemists to understand how structure might relate to activity in order to design even more active molecules. Computational chemistry came of age in the pharmaceutical industry, focused initially on methods of obtaining structure activity relationships (SAR) between series of active molecules, and then
evolving into de novo rational design efforts using structural information about the target molecule. The explosion of molecular biology went hand-in-hand with this evolution of rational design and mechanistic approaches by providing the means to obtain sufficient quantities of target proteins to both enable high volume assay as well as structural studies on otherwise inaccessible target molecules.

The extraordinary growth of combinatorial chemistry beginning some five years ago has led to yet another revolution, focused directly on the approach used by the medicinal chemist to generate lead compounds and to optimize their activity. However, the impact of combinatorial chemistry does not end with the successful discovery of therapeutic agents. It is now entering another phase even before its impact on lead generation and optimization has been fully developed or implemented by most companies, and even though the solid phase chemistry for non-peptide synthesis remains largely undeveloped. This new role of combinatorial chemistry is being spawned by the rush of genomics information which will produce an incredible number of new potential targets each year, more than overwhelming the capacity of the pharmaceutical industry. What is crucial is no longer the sequencing of the human and other genomes, but instead the identification of targets and their rapid evaluation to enable the industry to identify those with the greatest potential for impact on disease. To-date the use of transgenic experiments, and to a lesser extent, the use of antibodies, have served as the mainstay for determining the role of new, or even somewhat older target proteins in disease. However, combinatorial chemistry in the form of antibody display libraries, peptide display libraries, and synthetic peptide combinatorial libraries offer state-of-the-art tools to the pharmaceutical industry which can be exploited to identify antagonists/inhibitors and agonists of new target molecules which can in turn be used in vivo to validate the potential of exploiting a new target to address specific diseases. With time, non-peptide combinatorial chemistry may similarly be used to address this need, but currently such libraries are limited by the state of synthetic chemistry knowledge. While the potential for target validation is just now beginning to be exploited, results reviewed here clearly indicate that screening synthetic peptide libraries can produce leads for a large variety of targets of therapeutic interest, provide extensive databases of SAR to the medicinal chemist, and when libraries are designed based on the initial leads or other information, provide a valuable tool to rapidly optimize the activity of molecules. The envelope of peptide chemistry has been pushed to enable the reliable, high-purity synthesis of large numbers of peptides containing available proteinogenic and nonproteinogenic amino acids, amines, and carboxylic acids as linear, cyclic, and branched structures in order to produce the greatest diversity among library components and between libraries.

This chapter focuses on synthetic peptide/peptidomimetic combinatorial library synthesis, lead generation, and optimization. Numerous reviews have discussed the various formats used to synthesize combinatorial libraries: the
There is no universally accepted definition of combinatorial chemistry. For the purpose of this chapter it is defined to encompass the synthesis of compounds from sets of subunit and chemical reactions used in one or more reaction steps. This definition includes both solution and solid-phase synthetic reactions, the synthesis of compounds through several sequential reaction steps in which the same or different sets of subunits and chemical reactions are used, as well as the reaction of multiple subunits in one reaction step to form multicomponent compounds. The compatibility of each building block with each chemical reaction, and the product formed must be known and validated for the process to give useful libraries of compounds. While a distinction has been made in the past by some investigators between oligomer compounds and other multicomponent compounds, there is no difference between a linear or a branching synthetic scheme. Thus, just as in polymer chemistry where there may or may not be cross-linking and branching, so in oligomer chemistry where the subunits or monomers making up the final compound are different rather than the same (thus distinguishing the compounds from polymers of the same subunits) the oligomeric compounds may be linear or branched.
There are several basic strategies to the design, synthesis, and screening of combinatorial libraries. To simplify this review the different strategies are explained below, and the terminology that is used throughout this review is established. Multiple synthesis covers any method in which discrete compounds are synthesized simultaneously to create a library of isolated compounds whose identity is known from the synthesis scheme. Iterative synthesis/screening involves the synthesis of compounds in such a manner that a mixture results that is not directly resolvable to determine the identity of discrete active compounds, but that instead is resolvable to determine the identity of a specific residue(s) in any mixture that shows activity when assayed. A new set of mixtures is then synthesized based on this information and assayed, and the identity of the next specific residue(s) determined. The iterative process is continued until the identity of a complete, active molecule is determined. This process has several characteristics. First, the pool size diminishes as the iterations proceed, ending at the last step with the synthesis and testing of individual compounds. Second, the relationship of the last residues to be defined depends on the selection of the residues in the initial iterative steps, so there is an algorithmic relationship between the first and last components of the compound to be identified. Methods of deconvolution fall within the iteration definition and will be considered as a form of iterative synthesis/screening.

Positional scanning is a method of synthesizing several mixtures of compounds such that in each specific residues can be defined. The whole set of mixtures enables the investigator to identify every active residue of a virtual compound without actually having synthesized and tested that compound as a pure entity, only presumably as a component of each mixture. This process does not involve any iterative synthesis and assay steps. Furthermore, unlike the iterative process, there is no dependence of the identification of any one residue on the identity of any other residue within the active molecule. The actual synthesis of the derived active compound(s) must be performed separately, ideally as a multiple synthesis effort.

One-compound-one-bead postassay identification design (PAID) approaches (sequencable/encoded) covers a spectrum of approaches that share a synthesis scheme that assures that each resin particle contains only one structural compound. This is efficiently and practically achieved by applying the split-and-mix method of synthesis introduced by Furka and independently by others to assure that every amino acid couples in the intended equimolar ratio. This approach has the unique requirement for investigators to maintain the relationship between assay result and the bead on which the compound was synthesized so that they can utilize information on the solid-phase particle to identify the active compound or components of the active compound. It is principally in the method of identification that variations of this approach have been introduced. In some instances the compound itself can be directly sequenced or otherwise identified by analytical methods. In other cases the synthesis is carried out in such a manner that a coding molecule is also
synthesized on the solid-phase particle, and it is the coding molecule that is used to identify the active compound. The furthest afield approach introduced to date is the use of freely mixable capsules containing both resin and a microchip which can be read by radio-frequency methods. The basis for split-and-mix libraries lies in the freely mixable character of the support, whether it is a single resin bead or a collection of resin beads in some capsule or a tube of gel. While the one-compound-one-bead postassay identification approach to library synthesis absolutely depends on use of the split-and-mix synthesis process, iterative approaches can also use the split-and-mix method of synthesis.

The concept of the iterative synthesis of mixtures, positional scanning, and efficient screening in solution of one-compound-one-bead libraries raises the issue of synthesizing and testing mixtures of compounds rather than pure, highly characterized compounds. Scientists are accustomed to testing known compounds individually so as to minimize the possibility that some unknown factor or interfering substance may influence the observations being made. In screening large numbers of compounds, and particularly in combinatorial chemistry for the synthesis of large numbers of compounds, there is frequently a great advantage to synthesizing and assaying mixtures or pools of compounds. The immediate question is what risk does this entail, and what pool sizes can be used. The absolute answer must be judged on a case-by-case, experimental basis. However, as seen from all the examples given below, there does not seem to be an adverse effect of employing mixtures of compounds. Direct examination showed that pooling of hundreds of compounds did not mask positive compounds.

Analogs of the RGD peptide sequence were synthesized, pooled, and screened for binding to the gpIIb/IIIa platelet receptor. The library synthesized was of the form YGRGY_{X,X}, where Y_{1} consisted of the randomization of S, D, R, H, E, while 19 L-amino acids were randomized in the X_{2} position (omitting isoleucine) and 20 L-amino acids were randomized in the X_{3} position, creating a library of 1900 species. The library compounds were cleaved from the resin, pooled according to the residue incorporated at Y_{1}, and assayed. The YGRGD X_{2}X_{3} pool inhibited the binding of soluble gpIIb/IIIa receptor to fibrinogen-coated plates greater than any of the other pools, exhibiting similar activity assayed as a pool of 360 compounds as did control peptide, RGDS. The identification of YGRGD X_{2}X_{3} was predictable from previously described SAR, and the authors did not define X_{2}X_{3}; the focus of this study was to examine the effect of mixtures on assay results. In this study, assaying compounds as mixtures (of 360 closely related compounds) did not negatively influence the binding data and the ability to detect an active compound. The examples cited below all suggest that screening mixtures of compounds is a powerful approach for generating important information about the structure of active compounds, and that to avoid the use of pooling, when otherwise appropriate, is to unnecessarily lessen the amount of information that can be obtained for a particular system.
COMBINATORIAL SYNTHESIS OF PEPTIDES

Choice of Solid Support

Beaded polymer used for library synthesis has to fulfill certain criteria depending on the synthetic and screening strategy. For libraries based on the one-bead-one-compound approach, the size and substitution homogeneity is very important: To be able to evaluate the biological signal created by peptide released from a single bead one must be sure that the amount of compound released from each bead is uniform. Also important is the resin resistance to the formation of clusters (resin stickiness): Clusters will prevent the statistical free redistribution of resin beads and thus substantially lower the number of structures created. The ability of the resin to swell in both organic and aqueous media is especially important when target binding to the beads is used as the criteria for positive bead identification.

Polyacrylamide beads fulfill most of the above-mentioned criteria and was actually the resin used for evaluating technology feasibility. However, polyacrylamide has been replaced by TentaGel, polyoxyethylene-grafted polystyrene, which has become the resin of choice for both peptide and organic solid-phase library synthesis. The PEG-PS resin of Millipore (Perseptive Biosystems today has similar composition and properties, but differs in the placement of the chemically reactive group (amino group) in relation to the polystyrene matrix. TentaGel has the functionalizable group at the end of the polyoxyethylene chains, far from the hydrophobic polystyrene chain, a feature that is especially important for compound display on the bead surface for the bead-binding assay. PEG-PS has the functional group next to the polystyrene chain such that the polyoxyethylene chain does not serve as a linker connecting the synthetic compound with the polymer, but rather as modifier of polymer properties.

Alternative carriers to classical resin beads have been tested. Polyacrylamide-grafted polypropylene pins were used for the synthesis of the first library. This type of support was shown to be very useful in multiple-peptide synthesis. It was adapted to larger scale by the application of crowns attached to the pins, and is also useful for nonpeptide synthesis.

Paper is a good support for multiple (SPOT) synthesis of thousands of peptides or for the synthesis of libraries. Synthesis on paper is performed by spotting the solution of protected amino acids onto the functionalized cellulose paper in the presence of activating reagent. In this case, the reaction vessel is the carrier itself, liquid manipulation during the synthesis (usual shaking in the case of solid-phase synthesis) is eliminated, and the reaction is driven by the diffusion of liquid in the carrier. This principle of internal volume synthesis was tested using polymeric carriers on a multiple synthesizer utilizing centrifugation for liquid elimination and was found comparable, if not better, than the classical arrangement of solid-phase peptide synthesis. Cotton, the purest form of cellulose, was found to be a convenient solid-phase support, especially for multiple synthesis or library generation.
One specific feature of membrane, sheet, or thread-like carriers is their divisibility. This feature can be used for the synthesis of libraries with a nonstatistical or forced distribution of library members. The synthesis of this library starts with \( n \) pieces of the carrier, which are coupled with \( n \) different building blocks (1st randomization). Each of the \( n \) pieces is then divided into \( m \) parts and these smaller parts are distributed into \( m \) reaction vessels in which \( m \) reactions are performed (second randomization). The process can be repeated as many times as permitted by the handleability of polymer particles. The result of this process is a library of \( (n)(m) \cdots = X \) compounds on \( X \) polymeric particles, where no compound is missing and none is represented more than once.

**Synthetic Issues**

Coupling in peptide library synthesis is performed in the usual way—in a battery of bubblers, closed plastic vials, polypropylene syringes, or in tea bags—and standard coupling reagents are used. Detailed descriptions of library synthesis procedures have been published. Difficult couplings can be forced to completion by the application of more reactive coupling reagents, such as TFFA. However, due to the ease of handling and simplicity in deprotection performed in multiple vessels in parallel, the Fmoc protecting group is favored over the Boc group.

The techniques used to monitor coupling reactions required adaptation. In split-and-mix synthesis, beads within each reaction mixture contain different peptide sequences, and therefore coupling kinetics for each bead may be different. Since we are interested in knowing not whether the coupling is complete on average, but rather whether coupling is complete on each bead in the sample, it is advisable to follow the reaction at the level of individual beads. This can be achieved by nondestructive methods such as bromphenol blue monitoring. An example of monitoring individual beads is shown in Figure 3.1. Coupling was complete for the majority of beads (clear beads), but coupling on several beads is not complete (colored beads). The classical ninhydrin test did not reveal any problems at this coupling stage.

Equimolarity of incorporation of amino acids into the growing peptides of a library can be achieved in several ways. The most reliable is split synthesis, which was designed for the purpose of generating equimolar peptide mixtures. Equimolarity, however, can be achieved only in the cases when the number of particles used for the synthesis is substantially larger than the number of synthesized peptides (1,000,000 compounds cannot be synthesized on 100,000 beads). Split synthesis results in a collection of polymeric particles, each containing individual sequences, and was the basis for the development of the one-bead—one-compound library technique. The split method, however, is inconvenient in the case of iterative libraries containing fixed positions inside of the peptide sequence. In this case the synthesis of a library with 20 amino acids in one fixed position and 20 amino acids in positions requiring equimolar
incorporation of amino acids would require the use of 400 reaction vessels. An alternative is the use of a mixture of amino acids in which the ratio is adjusted according to the reactivity of the particular protected building block; or double or triple coupling of subequimolar (0.8 molar) amount of equimolar mixtures of amino acids. With either approach, the library would contain an equimolar representation of each peptide, though each bead would have a collection of peptides synthesized onto it (again with equimolar representation).

Synthesis of libraries can be performed manually or by automation. The distribution of the resin is achieved by volume distribution of either a homogeneous (nonsedimenting) suspension of beads in isopycnic solution or a stirred suspension (Saneii et al., 1993). The third design achieves distribution through sedimentation of the suspension in a symmetrical distribution vessel (Figure 3.2).

Libraries immobilized on glass slides have been synthesized by the application of photolithographical techniques. The identity of positively reacting molecules is known from the position on the glass surface (see Chapter 4). The solid-phase synthetic principle is also used in the case of liquid-phase combinatorial synthesis. Synthesis is performed on soluble polymer (functionalized polyoxymethylene), which is then precipitated for removal of excess reagents. Biological screening can be performed with compounds directly attached to the soluble polymer.
Figure 3.2. Redistribution of resin: The redistribution of resin into multiple vessels through the uniform settling of resin suspended in liquid media. The resin is thoroughly mixed and then allowed to settle. The design of the settling apparatus direct equal portions of the resin into the reaction vessels for the next step of synthesis.
Structure Determination by Nonencoded Means

Structure determination of the active component of a mixture is not an issue in the techniques utilizing iterative synthesis or positional scanning, the technique in which a certain feature of the library molecules is kept constant and the rest of the molecule structure is randomized. Determining the activity of variously defined mixtures defines the synthesis of the next generation library (or individual compound array in the case of positional scanning), and the active structure is defined by a synthetic algorithm. Part of the material used in each step of the synthesis of an iterative library can be saved and used for the synthesis of subsequent library generations. This principle, called recursive deconvolution, is attractive for laboratories evaluating one type of combinatorial library. It is not practical when hundreds of different libraries containing up to 100 building blocks in each randomized position are used, since one library composed of 4 consecutively connected building blocks, each position being randomized by 100 different blocks, would require storage of 1,010,100 resin aliquots.

A technique allowing the rapid determination of the active compound from a mixture is based on an orthogonal mixture principle. In this case two (or more) sets of mixtures are generated in such a way that individual compounds are present in two (or more) different mixtures in identical concentration. The biological activity of two orthogonal mixtures defines the active compound common for both mixtures. The disadvantage of this method is the complexity of the synthesis of the orthogonal libraries (Figure 3.3), but robotic synthesis of the library addresses this problem. Conceptually similar was a library design utilizing enriched and depleted mixtures of amino acids in the construction of library mixtures.

Library Formats

Challenging syntheses are the norm in combinatorial peptide chemistry, not the exception. For example, some biological targets require a free carboxy terminus (or both N- and C-termini free) to be displayed from amino to carboxy terminus. This is technically problematic due to significant problems with racemization in every step. This problem was solved by variously synthesizing a cyclic peptide containing a cleavable linker in its cyclic structure such that cleavage of the intramolecular linker exposed the free terminus of the molecule (Figure 3.4).

Screening one-bead-one-compound libraries for activity in solution requires multiple release of an equimolar quantity of compound from individual beads (vide infra). This can be achieved by attaching the compound to the bead via a linker, allowing release in two independent steps or by attaching the compound onto a mixture of linkers cleavable under different conditions. The first approach has the advantage of not decreasing the amount of released compounds, since the amount of compound on each bead is multiplied by the
Figure 3.3. Orthogonal Deconvolution of Mixtures: Synthesis of two sets of libraries using orthogonal overlapping building block subsets permits the direct identification of an active compound from the assay of both sets without resynthesis. Only one compound is common between the active sets. In this example mixtures contained 125 compounds each. Each set contained 125 sublibraries (as depicted for set A and Set B) which were assayed (total of 250 assays) in order to screen and identify the active compound from among the total 15,625 compounds.
Figure 3.4. Synthetic method for the solid phase synthesis and display of a free carboxy terminus of peptides remaining bound to solid phase particle during assay. The approach depends upon the synthesis of a peptide on a branched linker containing a protected arm, followed by deprotection of the linker arm and cyclization back onto the linker, and then cleavage of the original linker arm on which the linear form of the peptide had been synthesized. The peptide remains bound to the resin (indicated by the "P") through the cyclizing peptide-linker bond.

branching of the linker to accommodate each step of release (the release mechanism is illustrated in Figure 3.5). An alternative method of tiered release is the kinetic release realized by either timed exposure of a relatively stable (benzhydrylamine) attachment to mild cleaving conditions (vapors of trifluoroacetic acid), or exposure of photolytically cleavable attachment to light. Various approaches to stepwise release of the compounds from polymeric carrier have been reviewed.

The principle of a one-bead-one-compound library can be combined with the multiple-defined positional scanning concept to address the problem of generation and screening incomplete libraries. A library of hexapeptides composed from 20 amino acids would contain 64,000 compounds. However, it is probable that not all six amino acids in any one peptide with significant affinity for the biological target are essential for binding. Some of the amino acids in the sequence can be easily replaced without significant loss of activity, whereas several critical amino acids cannot usually be replaced by a substitute. The task is therefore to identify the critical amino acids, or the motif required for binding. If we define the motif as an arrangement of three amino acids, not necessarily contiguous, we can create 20 different positional arrangements
Figure 3.5. Chemistry and method of release of identical copies of a compound from a doubly cleavable branched linker. Branching of the linker enables two equivalents of peptide and one equivalent of coding molecule to be synthesized on the resin (indicated by the "TG") for every equivalent of resin loading capacity. The linker used permits one copy of peptide to be released by acid, removed and tested, and then the second copy of peptide can be released at neutral pH, removed and tested, and finally the identity of peptide can be determined by the coding molecule left on the resin bead. The released peptides have the identical chemical memory (hydroxyl rather than a c-terminal carboxyl group) of the
Figure 3.6. Method of synthesis of a "library of libraries": Each solid phase support contain a mixture of compounds representing a library of compounds ranging in length from 3 to 15 residues to evaluate all 3-residue pharmacophores. While it is not necessary for the 3-residue pharmacophore to be composed of contiguous amino acid residues, in peptides greater than six residues in length pharmacophores containing residues separated by more than three residue uni positions (e.g. the pharmacophore of amino acid 1, 5, and 9 of a nonapeptide) will not be represented.

of those three amino acids in the framework of the hexapeptide. In the case of 20 amino acids, each of these positional arrangements contain 8000 individual compositional motifs. Therefore, the complete library of tripeptide motifs in a hexapeptide framework, a "library of libraries," is composed of 160,000 species of motifs. This is a substantial reduction in complexity compared to the 64,000,000 individual compounds in a hexapeptide library.

The synthesis of a library of libraries requires splitting the resin into several aliquots—in the case of the above-mentioned libraries of hexapeptides, into up to six aliquots. In three aliquots a mixture of amino acids is coupled and in three aliquots the position is randomized between the 20 different amino acids during parallel coupling as in the standard one-bead-one-compound synthetic method. In total, 252 couplings are performed in the synthesis of this library (for the scheme see reference 61).

The scheme of synthesis of an alternative library with variable length is depicted in Figure 3.6. At the beginning of synthesis, and after each randomiza-
tion step, one-quarter of the resin is separated and the mixture of amino acids is coupled to the remaining part. After this coupling, one-third of the resin is separated and the remainder undergoes coupling with the mixture of amino acids. The next coupling is performed with half of the resin from the previous coupling. All portions of the resin are then combined and a randomization is performed. Synthesis of a library of libraries with a three-amino acid motif, by this method, consists of three randomization steps and four stages of multiple couplings of amino acid mixtures. As a result, each solid-phase particle of the library is subjected to three mandatory randomization steps and as many as 12 acylations with the mixture of amino acids. This library, containing peptides of lengths from 3 to 15 residues, consists of 256 positional motif sublibraries. Among sublibraries of up to hexapeptides, all positional motifs are presented. However, because this synthetic scheme does not allow more than three successive acylations with the amino acid mixture, motifs in which pharmacophore positions are separated by more than three adjacent structural unit positions are not represented.

A significant increase in the diversity of peptide libraries can be achieved by using not only alpha but also other amino acids in the construction of the peptidic chain. A peptide backbone can serve as the scaffold onto which a variety of building blocks can be attached via coupling to trifunctional amino acids (aminoglycine, diaminopropionic acid, diaminobutyric acid, ornithine, lysine, iminodiacetic acid, aspartic acid, glutamic acid, serine, hydroxyproline, cysteine, etc.). Iminodiacetic acid is a convenient structural unit allowing construction of peptide-like libraries. The scheme for its application to library construction is given in Figure 3.7.

Attachment of carboxylic acids onto the free amino groups was the basis for the construction of an alpha, beta, gamma library in which both the backbone and the side-chain arrangement was randomized. On the border of peptidic and nonpeptidic libraries are the peptoids, or NSGs (N-substituted glycine peptides). Libraries of peptoids were constructed either by coupling individual N-substituted glycines or by a submonomer approach, in which bromoacetic acid was coupled to the amino acid group of the growing peptoid chain, and primary amines were used to displace bromine in nucleophilic substitution, forming thus another N-substituted glycine in the chain (see Chapter 6).

The linear arrangement of building blocks in peptide-like fashion was the basis for several library designs. Because these designs produce nonpeptidic structures, they are not discussed here in detail.

Cyclization is a generally accepted method for decreasing the conformational flexibility of peptides. This restriction is expected to provide more potent ligands, as in the case of disulfide cyclic libraries containing ligand for the IIbIIIa receptor. Cyclization can change the preference of the biological receptor for ligands, as shown in the study of streptavidin binders from cyclic libraries of various sizes. A cyclic peptide composed of lysines and glutamic acid was used as a template onto which various carboxylic acids were...
Figure 3.7. Introduction of side chain diversity: Use of a trifunctional amino acid, such as the depicted iminodiacetic acid, as a scaffold on which to synthesize diverse side chains while extending the peptide backbone the desired number of residues. Solid phase is represented by the "TG".
Figure 3.8. Synthesis of a cyclic-turn mimic on solid phase (indicated by “P”): The low molecular weight mimic displays side chains within the same spatial arrangement as the tertiary-turn structure of proteins.

attached. Cyclic peptide libraries were studied by Spatola et al., and an optimal strategy for their synthesis was devised. Ellman et al., synthesized a β-turn mimetic library with the same goal (Figure 3.8). Conformational restriction has also been achieved by constructing libraries with a bias toward α-helical conformation. One such library was synthesized by randomizing four positions in the sequence of the amphipathic helix (YKLLKLLKKLKLK) on either the hydrophobic or hydrophilic sides of the molecule, from which peptides with increased antimicrobial activity were identified.

The combination of peptide structure with nonpeptidic elements has been used for library design. Potent and specific zinc endopeptidase inhibitors were identified in a library of peptides with the amino terminus modified by the Z-Phe(PO₃CH₂) peptidomimetic group. A similar approach was used in a Pfizer study of the endothelin antagonist developed by Fugisawa. N-terminal substitution was kept intact and all amino acids were randomized by an array of natural and unnatural α- and non-α-amino acids. Promising leads were generated from a library constructed using acylation of the peptide chain by an array of three building blocks containing amino and carboxyl functionality and capped by a set of carboxylic acids. The peptide chain in this case was used as the biasing element targeting the binding pocket of the Src SH3 domain. Vinylogous sulfonyl peptide synthesis was developed and libraries were used for the studies of synthetic receptors. Boc protected vinylogous sulfonyl chlorides coupling in dichloromethane catalyzed by dimethylaminopyridine monitored by bromophenol blue (for proper base[DBU] excess during the reaction) was used in the steps of library synthesis. Monitoring of the
Figure 3.9. Incorporation of a target-specific (e.g. HIV protease) pharmacophore (diol motif) as the basic scaffold of a peptide library synthesized on solid phase (indicated by "P") followed by cleavage and testing of the free peptides.
reaction was found to be critical. Using an excess of the base destroyed the sulfonyl chloride, and even an excess of sulfonyl chloride did not result in complete amino group modification. Libraries of synthetic receptors were generated by the combinatorial synthesis of peptides on scaffold molecules such as a macrocyclic tetramine or steroid molecule.

Palladium-mediated macrocyclization was used in the synthesis of cyclic libraries. The carboxy terminal lysine side chain was acylated by acrylic acid, and the amino terminal group of the linear peptide was acylated by iodobenzoic acid. Pd(O)-mediated cyclization provided clean product in high yield. The synthetic scheme of C2 symmetric inhibitors of HIV protease (Figure 3.9) serves as an example of solid-phase synthesis utilizing a specific feature of the target molecule (in this case the diol structure) to simultaneously protect the functional group critical to the function of the constructed molecules and at the same time use this protection as the attachment to the solid support.

LEAD GENERATION AND OPTIMIZATION

Drug discovery is composed of two major elements: lead generation and lead optimization. The research reviewed in this section demonstrates that combinatorial peptide chemistry is a powerful tool for generating leads using generic libraries designed on general rather than target-specific principles. Combinatorial peptide chemistry is also a powerful tool for the optimization of leads through the synthesis and screening of optimization libraries designed based on a known lead or SAR. It is particularly powerful when combined with traditional medicinal chemistry and molecular modeling, though published examples of this are just now beginning to appear. Table 3.1 lists the targets for which leads have been identified and/or optimized from synthetic combinatorial peptide libraries. This list covers a variety of classes of targets. In some cases, investigators have pursued these targets as a means of validating a new method or the design of a combinatorial library. In other cases, information has been sought about the target and its chemical selectivity. In still other studies, the objective has been to identify or optimize a lead that can be used to modify the activity mediated by a specific target. Details of how each class of target has been pursued are discussed in the following sections.

Identification of Antibody Binding Domains and Putative Peptide Antigens

The identification of peptides that bind specifically to antibodies can be used to define the immunizing epitope within the native antigen. In addition, compounds can be identified that bind and elicit the same function as the native antigen, but that are structurally different, constituting a mimeotope—a mimic of the native antigen epitope. In either case there is the potential for identifying useful vaccines from synthetic combinatorial peptide libraries. There is also the potential to identify peptides that bind a portion of the eCDR region but
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<td>Pin syn/solution assay, biased library</td>
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<td>Anti-Hemaglutin mAb</td>
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<td>Novel binder</td>
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<td>Vancomycin</td>
<td>Novel binder</td>
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<td>Autophosphorylation of library cpds, LCMS detection, biased library</td>
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<td>Acetylcholinesterase</td>
<td>Inhibitor SAR</td>
<td>Syn &amp; spot on TLC plate, novel detection, biased library</td>
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<td>Pins, biased library</td>
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<td>Antagonist SAR</td>
<td>Pin syn/solution assay, biased library</td>
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<td>Novel antagonist, SAR, optimize activity</td>
<td>Microplate syn, novel information rich library</td>
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*Split and Mix, One-Compound-One-Bead, Postassay Identification Format Libraries*

Anti-β-endorphin mAb | ID epitope | New method/on bead | 9 |
Anti-β-endorphin mAb | ID epitope | Double release | 55 |
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<tr>
<th>Target Molecule</th>
<th>Scientific Accomplishment</th>
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<tr>
<td>Anti-β-endorphin mAb</td>
<td>ID epitope</td>
<td>On-bead/radiolabel</td>
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<td>Anti-β-endorphin mAb</td>
<td>ID epitope</td>
<td>On-bead/coded</td>
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<tr>
<td>Anti-cMYC mAb</td>
<td>ID epitope</td>
<td>On-bead/coded</td>
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<tr>
<td>Anti-gp120 mAb</td>
<td>ID epitope</td>
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<td>ID epitope/vaccine</td>
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<td>B-cell surface idiotype</td>
<td>ID epitope</td>
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<td>105, 106</td>
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<tr>
<td>MHC Class I (HLA-A2 * B7) Strepavidin/avidin</td>
<td>ID binding motif</td>
<td>On-bead</td>
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<tr>
<td>Neural neurophysin</td>
<td>Novel/specificity</td>
<td>On-bead</td>
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<tr>
<td>Src, PI-3-kinase SH3</td>
<td>Novel/putative substrates from database search &amp; ligand/SH3 NMR structure determined</td>
<td>On-bead</td>
<td>110</td>
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<tr>
<td>c-AMP dep. kinase</td>
<td>Specificity of substrates</td>
<td>On-bead phosphor-ylation/biased library</td>
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<tr>
<td>Src kinase</td>
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<td>On-bead phosphor-ylation</td>
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<tr>
<td>Hepatitis A 3C endoproteinase</td>
<td>Specificity of substrates</td>
<td>On-bead/novel sequencing method, biased library</td>
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<tr>
<td>Stromelysin/collagenase</td>
<td>Novel/specificity of substrates</td>
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<tr>
<td>Endopeptidase subtilisin Carlsberg</td>
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<td>Edopeptidase subtilisin Carlsberg</td>
<td>Novel inhibitors</td>
<td>On-bead/novel indirect fluorescence detection method</td>
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<tr>
<td>Protein tyrosine phosphatase</td>
<td>Inhibitors, optimize specificity/activity</td>
<td>Solution, rf chip coding, biased library</td>
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<tr>
<td>Trypanosomal gPG-kinase</td>
<td>Trypanosom-specific inhibitors</td>
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<tr>
<td>Leukocyte elastase</td>
<td>Inhibitors, optimize specificity, activity</td>
<td>On-bead, biased library</td>
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<tr>
<td>Thrombin</td>
<td>Inhibitor SAR, novel lead/optimize activity</td>
<td>On-bead, biased library</td>
<td>125</td>
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<tr>
<td>Factor Xa</td>
<td>Novel inhibitor lead, SAR/optimize activity</td>
<td>On-bead (lead from generic library, optimize with biased library)</td>
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<tr>
<td>Elastase, subtilisin, trypsin, etc.</td>
<td>Inhibitor SAR</td>
<td>Solution/novel affinity column-sequencing method, biased cyclic scaffold library</td>
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<td>gp11b/llla receptor</td>
<td>Antagonist</td>
<td>On-bead &amp; double release</td>
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<tr>
<td>C5a receptor</td>
<td>Agonist–antagonist switch, SAR</td>
<td>Double release, biased library</td>
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<td>G-protein coupled receptors</td>
<td>Novel agonists &amp; antagonists, SAR</td>
<td>Double release, novel melanocyte functional assay, biased library</td>
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<tr>
<td>Antimicrobial</td>
<td>Growth inhibitors</td>
<td>Single release, agarose plaque forming assay</td>
<td>131</td>
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<tr>
<td>Indigo carmine</td>
<td>Small molecule</td>
<td>On-bead, direct color detection</td>
<td>132</td>
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### Iterative Synthesis and Screening Library Methods

| Anti-peptide mAb                        | ID epitope                                                                               | New method/solution         | 8          |
| Anti-peptide mAb                        | ID epitope                                                                               | Solution/novel pooling      | 51         |
| Anti-β-endorphin mAb                    | ID epitope                                                                               | Solution                    | 133        |
| Anti-β-endorphin mAb                    | ID epitope                                                                               | On-bead                     | 10         |
| Anti-β-endorphin mAb                    | ID epitope                                                                               | Liquid syn/screening        | 47         |
| Anti-TGFα mAb                           | ID epitope                                                                               | Solution, spot, double couple | 39         |
| Anti-gp120 mAb                          | ID epitope                                                                               | Robotics/affinity selection | 134        |
| Anti-LPS mAb                            | Discover novel epitope                                                                   | NMR friendly scaffold library | 135        |
### TABLE 3.1. (Continued)

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<tr>
<td>Anti-lysozyme mAb</td>
<td>Discontinuous epitope presentation</td>
<td>Rigid scaffold library</td>
<td>136</td>
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<tr>
<td>cAMP/cGMP kinases</td>
<td>Novel/substrate specificity</td>
<td>Solution</td>
<td>20</td>
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<tr>
<td>HIV-1 protease</td>
<td>Novel inhibitor/SAR, optimize activity</td>
<td>Solution, biased statin library</td>
<td>137</td>
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<tr>
<td>Trypsin</td>
<td>Novel inhibitor/SAR, optimize activity</td>
<td>Solution, lead generation/optimization</td>
<td>24, 138</td>
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<tr>
<td>Zinc endopeptidases</td>
<td>Optimize inhibitor selectivity/activity</td>
<td>Solution, biased library</td>
<td>116</td>
</tr>
<tr>
<td>Endothelin receptor</td>
<td>Antagonist SAR, optimize activity</td>
<td>Solution, biased substance P library</td>
<td>139, 140</td>
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<tr>
<td>Endothelin receptor</td>
<td>Antagonist SAR</td>
<td>Solution, biased antagonist FR-139317 library</td>
<td>77</td>
</tr>
<tr>
<td>μ-Opiate receptor</td>
<td>Novel agonists, SAR, optimize analgesic activity &amp; selectivity</td>
<td>Solution</td>
<td>141-3</td>
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<tr>
<td>Antimicrobial</td>
<td>Growth inhibitors</td>
<td>Solution</td>
<td>144, 145</td>
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| Positional Scanning Format Library Methods | | | |
|---------------------------------------------|--------|--------|
| Anti-peptide mAb | ID epitope | New method | 146, 147 |
| ACE and other enzymes | Substrate | Solution/novel seq. method | 148 |
| Chymotrypsin | Novel inhibitors | Tea bag, cyclic template library | 72 |
| IL-6 receptor | Novel antagonists, SAR | Solution, branched “polydentate” library | 149 |
| MHC class 1 | Novel ligands binding MHC-I, &/or sensitize target cells for cytolysis by T-cell | Solution | 150, 151 |
| RBC lysis | Inhibition of melittin mediated lysis | Solution | 152 |

*Unless noted, libraries used were generic in design. Biased libraries include overlapping, deletion, truncation, or biased amino acid selection based on sequence of native antigen or ligand.*
that do not elicit an antibody response, particularly when the native antigen binds through discontinuous epitopes or a conformational epitope. In addition, due to inherent tight binding and a wide variety of commercially available reagents and assays, antibody binding has been used as a method for validating new approaches in the application of synthetic combinatorial chemistry.

Multiple-Synthesis Libraries. In the first study published describing the multiple synthesis of peptides on fixed solid supports in a 96-well microplate array, the target used to validate the approach and demonstrate utility was an antibody. The pins, which had been dipped in reaction solutions to synthesize a library of known compounds, were dipped in assay medium containing a monoclonal antibody raised against the immunologically important coat protein (VPI) of foot-and-mouth disease virus to identify peptides that bound the antibody. The authors first synthesized 108 different overlapping hexapeptides required to cover the complete 213-amino acid sequence of VPI. From the results they identified GDLQVL as the epitope recognized by the antibody. They then synthesized a library of 120 different peptides in which all possible single-residue replacements were made using the 20 proteinogenic amino acids. This identified the two leucine residues as critical for antibody binding.

Another early study defining antibody–peptide recognition was reported by Houghten. He described the multiple synthesis of peptides by segregating resin in individual permeable containers (“tea bags”), then simultaneously coupling amino acids to the segregated resin of a number of such tea bags suspended in a variety of reaction vessels. Each tea bag was tracked through the synthetic scheme so that at the end of the process there was a library of compounds synthesized simultaneously in a one-compound–one-bag format, ready for screening after cleavage from the resin. Using this method 238 different 13-mer peptides were synthesized, reflecting single amino acid variations of a sequence from hemagglutinin protein (residues 98–110, YPYDVPDYASLRS), which has been used to raise a monoclonal antibody. The individual peptides cleaved from the bags of resin were preadsorbed to microtiter plate wells to constitute an ELISA screening assay, and the antibody binding to these peptides was monitored. Houghten demonstrated that with this approach the residues critical for binding could be identified as, and validating the tea-bag multiple-synthesis concept and utility.

Antibody binding was used to validate a multiple-synthesis approach in which peptides are synthesized as spots on commercially available peptide synthesis polypropylene membranes (Millipore, Bedford, MA) and then assayed either while attached to the membranes by an ELISA-type assay or following release from the membranes. A related synthesis method had been described using cellulose membranes. The membranes were mounted on a 96-well synthesis device. The library consisted of all possible octamers (1–8, 2–9, . . . , 269–276) derived from bovine myelin proteolipid protein (PLP, 276 amino acid peptide), which had been used as the antigen for producing
the target molecule anti-PLP antibody. Synthesis was complete in 16 h. The polypropylene sheet was then incubated with the anti-PLP antibody, and bound antibody was detected with protein A conjugated to alkaline phosphatase via a calorimetric enzyme assay. The dominant epitopes mapped by this approach corresponded to peptides spanning residues 38–46 and residues 195–205. The authors thus not only validated a novel method for multiple synthesis, but also identified the binding epitopes for this anti-PLP antibody.

Frank et al. used serum from humans that was immunopositive for a 58-amino acid sequence from human cytomegalovirus 36/40 K protein (anti-CMV) as a test system. They synthesized a library of 49 overlapping decapeptides on a sheet of cellulose paper, spotting the reaction mixtures as separate points of a grid to form an array of compounds. With the peptides still bound, the cellulose paper was incubated with the CMV positive serum. Positive spots were detected by identifying those to which antibody from the serum bound (using an anti-human β-galactosidase-conjugated second antibody). In this way the authors could map the binding epitope of individual human sera. This method was also used to characterize the epitope recognized by an anti-TGFα antibody, Tab2, by synthesizing a hexapeptide library spotted on cellulose of 1728 mixtures in the format XOOOOX, were X represents the positions where residues were randomized, defined by the spot location, and O represents positions where one of six amino acid mixtures was incorporated (A,P,G; D,E; H,K,R; N,Q,S,T; F,Y,W; I,L,V,M). A similar library was generated by phage display. The synthetic combinatorial library identified sequences that were native to TGFα (SHFNDC, \( K_d = 200 \text{ nM} \), and VSHFND, \( K_d = 80 \text{ nM} \) ) as well as a large number of other sequences with comparable affinity. The phage library identified the native HFND sequences (ASHFND, \( K_d = 60 \text{ nM} \); HFNDYL, \( K_d = 900 \text{ nM} \); and HFNDCL, \( K_d = 8000 \text{ nM} \) ), but, unlike the synthetic library, not the complete native sequence VSHFND or any novel sequences.

Stigler et al. characterized the binding epitope of the Fab fragment of an antibody (3D6) raised against the transmembrane protein gp41 of HIV-1 using a library of overlapping peptides, a random library, and molecular modeling. The overlapping peptide library identified CSKLICTTAVPW, which was used to synthesize a secondary library on cellulose sheets to generate a library of all possible L and D conformers (494 in all). This identified critical residues and this information, together with low-resolution X-ray of the peptide bound to the Fab fragment, enabled the authors to construct a model of the bound conformation of the peptide and interactive residues. This study provides a strategy for constructing models of protein–peptide complexes using peptide library results to generate empirical experimental data to augment computer modeling and low resolution crystallographic approaches.

A useful method for determining antibody binding epitopes was demonstrated using photochemistry and light-directed synthesis on a sheet of glass to create an array of overlapping epitopes against which to screen an antibody. The epitope of an antibody, C 32.39 anti-dynorphin-β-antibody, raised
against the opioid sequence YGGFLRRQFKVVT was identified by synthesizing a library of 1024 compounds consisting of the single amino acids as well as all di-, tri-, (etc.) peptides to cover all possible deletions (single, double, etc.) and truncations of, while not scrambling, the original sequence. This represents a very different strategy than simply the synthesis of overlapping sequences within the immunizing peptide. Fluorescently labeled antibody bound to the individual compounds displayed as an array on the glass surface was monitored, with the location on the glass giving the identity of each compound. Deletion sequences containing RQFKV (four in all) exhibited high fluorescence. From the truncation results, loss of T, especially T plus V, from the C-terminus resulted in a loss of activity, while binding increased with truncation of the N-terminus to the second R, again identifying the RQFKV(T) sequence as the epitope recognized by antibody. H-YGGFLRRQFKVVT-OH bound with an IC₅₀ of 0.58 nM, while Ac-RQFKVVT-OH bound with an IC₅₀ of 1.7 nM, and Ac-RQFKVVT-NH₂ bound with an IC₅₀ of 0.49 nM.

Numerous other studies have demonstrated the utility of synthetic peptide libraries in the mapping of antibody binding domains and the production of potent antigens. Pinilla et al. recently demonstrated that calcium independent antigens could be identified by screening a library against a calcium-dependent monoclonal antibody; the anti-FLAG antibody M1, which binds DYKDDDDK-NH₂ in the presence of calcium and is used for protein purification. The IC₅₀ of DYKDDDDK-NH₂ determined with and without calcium was 14 and 274 nM, respectively, while a peptide identified from a hexapeptide library, DYKAKE-NH₂, exhibited an IC₅₀ of 38 versus 3 nM in the presence and absence of calcium, respectively, and one identified from a decapeptide library, DYKEKELEDD-NH₂, exhibited an IC₅₀ of 19 versus 2 nM with and without calcium, respectively.

Experiments with an anti-gp120 antibody identified an equipotent sequence containing nonnatural residues from a highly biased decamer library based on the native sequence Ac-RAFHTTGRINH₂. The library, Ac-RA(X)HTTG(X)I(X)-NH₂, produced an equipotent submillimolar affinity compound, Ac-Rα-naphthylalanine)HTTG(R)I(L-norvaline)-NH₂, and related naphthylalanine containing analogs due to the pooling method used in screening.

The multipin scanning approach of combinatorial peptide synthesis was used to identify the minimal and optimal length of an endogenous sequence from HIV-1 gp160, which serves as the recognition domain for binding to the MHC class I D₄ complex of murine cytotoxic T lymphocytes (CTL) to elicit a specific CTL response (lysis of peptide-binding, D₄-expressing target cells). The approach consisted of synthesizing an array of either N-terminal and C-terminal truncations of the 15-mer sequence from the native protein (residues 315–329). The minimal sequence required for recognition based on this experiment consisted of an 8-mer, PGRAFTI, residues 320–327. In the second set of experiments the C-terminus was either extended or truncated, and for each C-terminus (331, 330, 329, 328), a set of six different N-terminally
truncated peptides (residues 318 to 323) was synthesized to create, for example, six 331 C-terminal peptides with an N-terminus of residue 318, 317, 316, 315, 314, or 313, respectively. Assaying this set of compounds determined that truncation beyond residue 320 resulted in loss of activity regardless of the C-terminal extension, confirming the minimum sequence, and identifying the most active peptides as the 9-mers PGRAFVTIG and GPGRAFVTI or the 10-mer RGPGRAFVTI (residues 318–327).

A corroborating study using a conventional approach also identified the 10-mer (residues 318–327) as more active than the native 15-mer. When the 10-mer peptide was incorporated into VAC recombinants and used to immunize mice, CTL lines obtained from the immunized mice exhibited specific priming by synthetic peptide comparable to that induced by wild-type, full-length gp160, demonstrating that peptides identified by synthetic combinatorial chemistry efforts can be used effectively as the basis for vaccine production.

**Split-and-Mix, One-Compound–One-Bead, Postassay Identification Format Libraries.** Antibody binding was used to validate the use of the split-and-mix approach to synthesizing and screening libraries in a one-compound–one-bead, postassay, directly sequenceable identification format. A library of peptides was synthesized on beads using the split-and-mix strategy to assure that each amino acid coupled regardless of its inherent reactivity and to assure that there would be only one species of compound on each resin particle (Figure 3.10). With the peptides still attached to the beads, the library was incubated with mouse monoclonal anti-β-endorphin. This is an antibody that recognizes a sequence in the native ligand (binding affinity, 17 nM). Antibody binding to beads was detected using a secondary antibody conjugated to an enzyme reporter (e.g., alkaline phosphatase), the positive beads were collected, and the peptide on each was sequenced to determine the identity of the individual compounds. The ligands identified from peptide libraries in which 19 of the proteinogenic amino acids were coupled at each position included YGGF(X) motifs, among them YGGFQ (15 nM affinity). The authors demonstrated that a positive bead could be treated to remove the target molecule and reassayed to confirm binding or to assess specificity. Furthermore, the bulk of the on-bead library remained after screening (minus any positive beads) and could be reused in additional screens.

A significant problem with the use of streptavidin to identify the bound biotinylated antibody is that streptavidin itself binds to many peptides, and hence can bind directly to beads (examples given below). To avoid this, two independent methods can be used to recognize the bound biotinylated antibody: streptavidin and an anti-mouse antibody. (Lam et al, 1995). Beads were first labeled with the streptavidin/alkaline phosphatase conjugate, colored (turquoise) with the substrate BICP, and selected. The selected beads were incubated with anti-mouse antibody conjugated with alkaline phosphatase, and colorized with BCIP plus NBT, turning positive beads dark purple.
Those beads remaining turquoise (not binding the anti-mouse antibody) were sequenced, and it was shown that the peptides were streptavidin binders. Those that turned dark purple proved to be YGG motif peptides, true anti-β-endorphin binders.

In a follow-up study the approach was further modified to permit the controlled double release and assay of the library peptides in solution. A pH-sensitive linker was employed that was stable in aqueous solution at acidic pH (4.5) but underwent a ring closure and cleavage to release compound (one-third of the total) from each bead at pH 7 in the form of...
XXXXX-NH-(CH$_2$)$_2$-OH.$^{56}$ Released in randomly generated pools of 500 beads per well, the mixtures were assayed in an ELISA-type assay to identify active pools. The beads corresponding to these active pools were then recovered and redistributed, 1–2 per well, and subjected to basic pH (0.2% NaOH) to cleave another third of the compound attached through an ester linkage, again releasing molecules with the structure XXXXX-NH-(CH$_2$)$_2$-OH. After assay and identification of the positive wells, the corresponding beads were recovered and sequenced to determine identity (using the remaining third of compound which had been synthesized onto a protected linker). The identified compounds were synthesized and their activity was confirmed. This follow-up study validated the double-release assay through the identification of the spiked control YGGFL (30 nM) as well as YGGFG (200 nM), YGVFG (1000 nM), and YGAFG (700 nM) from the assay of 500,000 library beads. The double-release assay is a powerful technique, now widely used by many laboratories.

In a related study the on-bead assay was conducted by incubating beads with radiolabeled target molecule, immobilizing the beads in a thin layer of agarose, identifying positive beads by autoradiography, and manually selecting beads for sequencing.$^{59}$ The target used was again an anti-β-endorphin monoclonal antibody. From a pentapeptide library the authors identified YGSFE, YLWFQ, and YGAFE, clearly related to the known YGGFL sequence, and in the case of YGAFE, nearly identical to a peptide found using the double-release assay described above. Like Lam et al.$^{9}$ the authors demonstrated that the recovered beads could be treated to dissociate the target and reassayed several times.

An on-bead method using a sequenceable coding molecule, rather than relying on sequencing of the active compound itself, was validated using the same antibody.$^{106}$ The assay was carried out using fluorescently labeled antibody and a fluorescence activated cell sorter to select the (positive) fluorescent beads. A heptapeptide library was synthesized on 10-mm macroporous beads, which permitted the target protein to permeate the bead and reach compound synthesized inside. The method was first validated using the known epitope RQFKVVT ($K_d = 0.5$ nM) synthesized onto beads. Two percent of the library beads stained above background and the bound fluorescent antibody could be competed by excess RQFKVVT, demonstrating specificity of binding. Peptides were identified from the library that, when synthesized with the threonine linker (T), ranged in binding affinity from a $K_d$ of 0.3 to 1400 nM. The sequence of the most potent ($K_d = 0.29$ nM) was TFRQFKV(T), a close analog of the native epitope RQFKVVT (T) ($K_d = 0.51$ nM).

An alternative coding method was validated using a monoclonal anti-c-MYC antibody as the target protein.$^{107}$ In this case the coding molecules consisted of a binary digital code arrangement of electrophoreses that could be separated and detected by electron capture gas chromatography (see Chapter 14). In this initial work the codes were attached to a fraction of the test compound, but the coding molecules can also be synthesized directly onto
the polystyrene bead. In digital coding the position and identity of each residue is represented by a three-“residue” bit code, where no residue is considered one bit of information, just as would be a specific coding residue. The library synthesized was a heptapeptide library of the format H₂N-XXXXXXEEDLGGGG-bead, containing D, E, I, K, L, Q, and S randomized at each position. It was known that the anti-c-MYC antibody 9E10 bound EQKLISEEDL, hence the selection of amino acids for the library. Using the approach described above (mixing library beads with the anti-c-MYC antibody and detecting those beads to which antibody bound using a secondary alkaline phosphatase conjugated antibody) the authors identified and sequenced the codes for 12 beads. Upon resynthesis and testing in solution the three most potent (IC₅₀ ~ 1 mM) were EQKLIS(EEDL), LQKLIS(EEDL), and QQKLIS(EEDL), all closely related to the known epitope, validating this coding methodology.

Screening using the anti-gp120 antibody was used to validate a laser desorption mass spectroscopic method of sequencing to determine the identity of compounds on positive beads. A synthetic hexapeptide library was synthesized in such a manner that at the end of each synthetic step a portion of the sequences were capped to prevent their complete synthesis in the remaining scheme. Thus, each bead contained the complete test peptide as well as capped truncations. The library beads were incubated with the anti-gp120 antibody, and positive beads were identified using a secondary antibody conjugated to alkaline phosphatase. Each positive bead was selected and its peptide contents cleaved and subjected to laser desorption mass spectroscopy. The spectrum contained the molecular ions for the complete compound plus all the possible capped analogs. The differences in molecular weight could be used to identify the composition of the parent peptide. This method can be applied to any molecular library, not just peptide libraries.

From the first study a family of peptides were identified, which provided a consensus binding epitope of (F/P)GRAF(Q/X)(F/X). This corresponded well with the peptide from the HIV gp120 protein used to produce the antibody originally, RIORPGRAFVTIGK (IC₅₀ = 11 nM), compared to an IC₅₀ = 50 nM for PGRAFQF. In the second study a pentapeptide library was synthesized, with and without acetylation, and screened, resulting in the identification of a XGRAF motif and an Ac-PG(R/X)AF(X/R)F motif. Ac-PGRAFQF exhibited an IC₅₀ of 525 nm. The authors also screened against streptavidin (see below), identifying the HPQ motif with and without acetylation.

Another example of identifying a vaccine from a synthetic combinatorial library has been published. Steward et al. synthesized an octamer library, screened on-bead with a monoclonal antibody raised against the F protein of measles virus (anti-MVF antibody F7-21). Six peptide mimotopes were identified, and when used to immunize mice, one (NIIRTKKQ), which cross-reacted with the antibody, inhibited measles virus plaque formation and acted as a vaccine to protect mice against fatal encephalitis induced by infection with measles.
Iterative Synthesis and Screening Library Methods. Hexapeptides were identified that bound a monoclonal antibody that recognized a 13-residue peptide (Ac-YPYDVPDYASLRS-NH₂) using an ELISA-type assay with the library peptides free in solution to identify antibody binding epitopes. A total of 324 different pools were synthesized with the format Ac-O₁O₂XXXX-NH₂ in which the first two positions of each were defined for each pool. Screening identified one pool containing Ac-DVXXXX-NH₂. Subsequently, 20 pools were synthesized with the format Ac-DVO₁XXX-NH₂ and screened to define O₁; the process iterated until each position had been defined. In the process of defining the last position a set of individual peptides rather than mixtures was synthesized and assayed, having the same O₁O₂ residues, but variable O₃ residues. Among them was the native sequence Ac-DVPDYA-NH₂ (IC₅₀ = 30 nM), which was also the most active. Corroborating studies using traditional approaches confirm this to be the antigenic determinant of the 13-residue peptide.

Using this approach the same group defined the binding epitope (STTS) of a monoclonal antibody raised against a surface antigen of hepatitis B, identifying the peptides Ac-STTSLI-NH₂ and Ac-STTSLM-NH₂ (IC₅₀ ~ 1.8 and 2.6 nM, respectively). Similar results were found using a positional scanning library approach (see below).

The success of the iterative approach, together with a novel pooling strategy, was also demonstrated using both an antiserum to a tetrapeptide as well as a monoclonal antibody to a 28-amino acid peptide. The antiserum used was raised against the peptide FMRF amide, but because the authors wished to exclude methionine, cysteine, tryptophane, threonine, and isoleucine from their library (reducing the number of amino acids used to 15) the ligand bound to microplates forming the basis of their antibody capture ELISA assay was FLRF. This strategy was somewhat unusual. The authors divided their amino acid mixtures into three groups, a(L,A,V,F,Y), b(G,S,P,D,E), and g(K,R,H,N,Q), with the individual amino acids present at different molar ratios designed to compensate for differences in reactivity. One strategy would have been to synthesize the library as 81 pools representing all possible positions within the tetramer and each of the three groups of amino acids. The authors instead synthesized a pool of compounds containing equal proportions of all 15 amino acids at each position (a 1:1:1 mixture of amino acid groups a:b:g), measured the activity of this pool (after the compounds were cleaved from the resin), and then compared this to at first four pools, which consisted of

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<td>Activity/Interpretation</td>
<td></td>
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<tr>
<td>1:1:1 a:b:g</td>
<td>1:1:1 a:b:g</td>
<td>1:1:1 a:b:g</td>
<td>1:1:1 a:b:g</td>
</tr>
</tbody>
</table>

| 2:1 b:g |
| 1:1:1 a:b:g |
| 1:1:1 a:b:g |
| 1:1:1 a:b:g |
| 1:1:1 a:b:g |
| 2:1 b:g |
| 1:1:1 a:b:g |
| 2:1 b:g |
| 1:1:1 a:b:g |
| 2:1 b:g |
| 1:1:1 a:b:g |
| 2:1 b:g |
| 1:1:1 a:b:g |
| 2:1 b:g |
Proportionate loss of activity was used to identify critical residues. The first round of screening results indicated a group residues were required at positions 1, 2, and 4 for activity, and g group residues were necessary at position 3. The next round was designed to identify triplets of amino acids from among these pools, using as the null control a, a, g, a, and four libraries:

```
<table>
<thead>
<tr>
<th>AA1</th>
<th>AA2</th>
<th>AA3</th>
<th>AA4</th>
<th>Activity/Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>a</td>
<td>g</td>
<td>a</td>
<td>null control</td>
</tr>
</tbody>
</table>
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The last round used the defined dipeptide mixtures as null control and the following four pools:

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<table>
<thead>
<tr>
<th>AA1</th>
<th>AA2</th>
<th>AA3</th>
<th>AA4</th>
<th>Activity/Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>F/Y</td>
<td>A/L</td>
<td>K/R</td>
<td>F/Y</td>
<td>null control</td>
</tr>
<tr>
<td>F</td>
<td>A</td>
<td>K/R</td>
<td>F/Y</td>
<td>=;F</td>
</tr>
<tr>
<td>F/Y</td>
<td>A</td>
<td>K/R</td>
<td>F/Y</td>
<td>&lt;;L</td>
</tr>
<tr>
<td>F/Y</td>
<td>A/L</td>
<td>K</td>
<td>F/Y</td>
<td>&lt;;R</td>
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<tr>
<td>F/Y</td>
<td>A/L</td>
<td>K/R</td>
<td>F</td>
<td>=;F</td>
</tr>
</tbody>
</table>
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Thus with three rounds of iteration, synthesizing 12 pools of peptides representing 50,625 peptides, the authors had defined the peptide FLRF.

Having shown that the approach worked, the authors then applied the method to the case of the monoclonal antibody, raised against the 28-amino acid peptide Ac-RTPALGPQAGIDTEIALEPDAPPDAC amide, and for which the epitope had not been defined. The difference was that a hexapeptide format was pursued, so that there were 6 pools in each round of iteration (18 pools synthesized, representing 16,777,216 peptides), but otherwise the strategy was identical. The result was the definition of RQVGH amide. The authors used this to identify the consensus region of the 28-mer to be PQAGID (bolded above), and confirmed this by synthesizing seven peptides and assessing their relative activity (shown in brackets) compared to the "native" peptide PQAGID [2.8]; RQVGH [1.2], QVGH [>750], RQVGH [128], EQVGH [14], PQVGH [0.08], RQAGHD [4], and RQVGD [17]. The loss of activity in the N- and C-terminal deletion analogs demonstrated specificity, and the ~35-fold improvement in activity of PQVGH over the native peptide PQAGID confirmed specificity and demonstrates the power of the approach for optimization.
In another study, peptides that bind anti-β-endorphin (3E7) were elucidated using iterative synthesis and screening of a series of hexamer libraries (Figure 3.11). Peptides were cleaved from resin as mixtures and assayed free in solution without separation in an ELISA-type format. The antigen peptide (β-endorphin) was adsorbed onto the surface of microplate wells, the peptide mixture plus antibody was added, incubated, and washed, and then bound antibody was detected. In the first stage 400 libraries were synthesized, each containing specified amino acids (O₁,O₂) in the first two positions, with the remainder (X) synthesized by splitting and mixing and cleaved as a complete mixture as regards the X positions (format O₁O₂XXX). Once the first two positions (A₁A₂) were defined, 20 libraries were synthesized, with the third position (O₃) identified in each (format A₁A₂O₃XX). This process continued until the complete peptide was defined, YGGFMT-NH₂ (IC₅₀ = 3.2 nM), a peptide identical to the first six residues of β-endorphin.

The most recent variation on the iterative approach is single-step deconvo-
Recursive deconvolution where automation is utilized to simplify the iterative process (Figure 3.12). In recursive deconvolution the library is split after each synthetic step into samples that are saved for subsequent synthesis and samples that are pooled and split in a many-compounds-one-well format (rather than the one-compound-one-bead format) for the next round of synthesis. Thus, for instance, at the end of the synthesis of a tripeptide library there would be multiple wells containing pools of compounds with the identity of the last amino acid unknown. Screening would identify the last amino acid, and then the investigators would take the previous saved plate of dipeptide mixtures, in which the second residue was known because of well location, synthesize onto these mixtures the identified third residue, and then screen to identify the second residue. Once identified, the investigators would go to the first round of saved single amino acid plates, synthesize the second and third residues onto each, and identify the active tripeptide. The authors synthesized a pentapeptide library, but with only four amino acids (G, L, F, Y) used at each position to create a library of 1024 members, including a peptide to bind the anti-β-endorphin antibody. This approach eliminates the need for
LEAD GENERATION AND OPTIMIZATION

tags, dependence on resin bead size to provide sufficient assay material, and some of the synthesis steps required in the iterative approach. The authors screened the library compounds in the anti-β-endorphin antibody ELISA assay with compounds attached to the beads to assess whether compounds in the mixture prevented the antibody from binding to the [Leu]enkephalin–BSA-coated surface. The authors identified the known peptide NH₂-YGGFL as well as two other peptides, NH₂-YGGFF and NH₂-YGGLL, validating their recursive deconvolution method.

An important issue in all the approaches where a mixture of amino acids is reacted in the same reaction vessel is the fact that some will couple at a faster rate than others, producing an unequal representation within the pool. The rates of reaction are somewhat inherent in the amino acid itself, but are also dependent on the sequence to which the amino acid is being coupled. Many investigators have tried to manage this problem by changing the ratio of amino acids within the mixture inversely according to their general reactivity, using a larger number of equivalents of the slower reacting amino acids to create a “kinetically adjusted” acylating mixture. However, another method has also been described. It had long been accepted in peptide chemistry that for some difficult couplings a process of double coupling was necessary to completely react all available sites on the solid phase. Double coupling means that after the first reaction is complete, the same reaction is repeated again to force coupling to all available reactive sites.

This approach was modified by Kramer et al. to assure that in a mixture there would be essentially an equimolar representation of all amino acids, regardless of the rate at which they reacted with the solid phase. They used a less than saturating mixture in the first coupling reaction, for example, 0.8 equivalents (relative to the number of reactive sites in the solid phase) of an equimolar mixture of amino acids. Thus in the first step the slowest reacting amino acid would be coupled to the solid support with close to the same efficiency as the fastest reacting amino acids. The result was an equimolar representation occupying 80% of the available coupling sites on the solid support at the end of the first reaction. The remaining reactive sites were coupled in the second reaction of the double-coupling protocol, so the fastest reacting amino acids coupled to a <20% excess over the slowest.

The authors validated this approach through the synthesis of libraries on solid-phase resin and on a cellulose sheet, using a variety of targets, among them the epitope definition of an anti-TGFα monoclonal antibody, Tab2. The iterative library design was XX₀₁O₂XX, where in this case the double coupling of a mixture of 19 amino acids was used at the X positions, and O₁ and O₂ were defined, creating pools of peptides in solution following cleavage from the solid support. A standard competitive ELISA was performed using TGFα bound to microplates and identifying those wells to which horseradish peroxidase labeled anti-TGFα did not bind in the presence of the particular peptide pool. The known epitope was HFND. From the library (containing 47,043,881 compounds in 361 pools) the authors identified XXHFXX, XXFNXX, and
XXNDXX, clearly peptide pools related to, and containing, the known epitope. When these pools were synthesized as spots in a 19 x 19, O, by O, array on a sheet of cellulose paper, the same families and conservative substitutions were identified: FN, HF, HY, ND, YF, YN, YY. Using two additional 19 x 19-array cellulose-bound libraries the authors identified the amino acids permitted on either side of the O,0, motif, synthesizing libraries of the structure X O, HF O,X, X O,FN O,X, and X O, ND O,X.

Positional Scanning Format Library Methods. A decapetide positional scanning library approach was evaluated using the same anti-Ac-YPYDVPDYASLRS-NH$_2$ antibody. Ten different positional libraries were synthesized, each containing 20 pools with a defined amino acid (O) at the specific position being investigated by that library (O,X,X,X,X,X,X, etc.). From this study 15 peptides were identified with comparable or better activity than the antigenic 13-mer (IC$_{50}$ = 6 nM), and all but one contained a consensus DYA sequence at positions 8, 9, and 10.

A derivative positional scanning method published by Wong et al. was also validated using antibody binding. The authors synthesized three libraries of the format Ac-O,O,XXXX-NH$_2$, Ac-XXO,O,XX-NH$_2$, and Ac-XXXXO,O,-NH$_2$, using only 10 "physicochemical representative" proteinogenic amino acids so that each library consisted of 100 dipeptide defined pools of 10,000 compounds each. Screened against a monoclonal antibody against the pili of Pseudomonas aeruginosa, the authors identified the best dipeptides, EQ, FL, and PK, corresponding to the known epitope DEQFIPK. They synthesized EQFIPK, and determined an IC$_{50}$ of 1.3 mM, validating the approach. However, when the 8 peptides containing single-amino acid substitutions of the residues representing amino acid classes (D for E; N for Q; W, Y for F; V, M for I; H, R for K) were synthesized and tested, only four exhibited activity similar to that of the lead sequence (IC$_{50}$ = 3.4-0.9 mM). This meant that while the representative algorithm was effective in enriching the active hit rate upon secondary synthesis, if the wrong representative residues had been selected no lead in the case of H or R for K or a less informative lead in the case of N for Q, or Y for F would have been discovered. Therefore, the authors propose that this approach is best used where it is necessary to simplify the synthetic process, such as synthesis and screening libraries of longer peptides (heptamer, octamer, etc.), trading off the risk of missing leads with the ability to actually synthesize and screen a complete library.

Using this approach Pinilla et al. defined the binding epitope (STTS) of a monoclonal antibody raised against a surface antigen of hepatitis B, identifying the peptide consensus Ac-S(S/T)T(P/S)(A/M)(H/M)-NH$_2$, from which the three most active peptides synthesized were Ac-STTSMM-NH$_2$ (IC$_{50}$ = 0.39 mM), Ac-STTSMM-NH$_2$ (IC$_{50}$ = 0.87 mM), and Ac-STTSAM-NH$_2$ (IC$_{50}$ = 2.8 mM). These results were essentially the same as those obtained by the iterative synthesis library described above.
**Liquid-Phase Combinatorial Libraries.** As combinatorial chemistry is extended from peptides to nonpeptides, synthesis issues become the major limitation. A method termed liquid-phase synthesis was demonstrated in which the compounds are synthesized onto a chemical phase (polyethylene glycol, MeO-PEG 5000) that can be conveniently crystallized or precipitated by solvent changes. Liquid-phase synthesis permits filtration purification and affords all the advantages of solid-phase synthesis when it comes to the separation of product and reactants, yet permits the reactions to be carried out in solution. This method was validated using anti-β-endorphin as the target and the recursive deconvolution strategy described above. From a pentapeptide library containing G, L, F, and Y, the authors identified YGGFL-MeO-PEG, YGGFF-MeO-PEG, YGGFY-MeO-PEG, and YGGFG-MeO-PEG using the same ELISA assay as described above, though in this case the pools of library compounds were in solution, though not cleaved from the PEG.

**Validation of Synthesis, Assay, and Compound Identification Methods.** The first description of a fully automated, 96-well, robotic, solid-phase library synthesizer was validated using an anti-gp 120 antibody. This system has been a model for many of the systems now in use and was based on the use of filtration microwell plates, enabling the removal of solutions without loss of the solid-phase support beads and the addition of solutions to the wells using a single- or multiple-head pipetting device. Individual compounds can be synthesized in each well or, by transferring the beads from wells into a mixing device and then redistributing them among the reaction wells, a library could be synthesized using either the split-and-mix one-compound-one-bead approach or any of the iterative, deconvolution, or positional scanning approaches.

In the example 12 individual peptides were synthesized and tested to validate the quality of synthesis. A library of format RAX,HTTGRIX, (19 amino acids randomized at each X position) was synthesized and assayed in pools of peptides released from the solid-phase supports using an ELISA assay. After addition of the X, amino acids the 19 pools (containing 19 peptides differing in X,) were kept separate so that the residue(s) in this position would be defined by the initial assay. The anti-gp120 antibody used had been raised against recombinant gp120, and the binding epitope mapped to a 10-mer sequence (RAFHTTGRII). The residues in boldface type had been shown to be important using alanine scans. Screening identified X, as F = W = Y > H > A = N. Each mixture was separated by affinity chromatography using the anti-gp120 antibody as the affinity support, and the retained fractions were sequenced to identify X, with the finding I > L > P. It would have also been possible to resynthesize the 19 peptides corresponding to each X, pool in order to identify X, but affinity selection as demonstrated in this work can be a powerful approach in certain circumstances.
Identification of Protein and DNA Binding Peptides

The binding to streptavidin has been used to validate new synthetic combinatorial methods. In general, either the direct binding of (alkaline phosphatase-conjugated) streptavidin/avidin to beads is monitored or biotin is bound to microtiter plates, test peptides are added free in solution, and the inhibition of streptavidin/avidin binding is monitored. While not of therapeutic interest, these studies are of scientific note, since they explore the binding of peptides to a protein that binds a nonpeptide (biotin) with very high affinity. Both L- and D-amino acid-containing peptides are bound by streptavidin and avidin. Other more biologically relevant protein-protein interacting systems have also been studied.

Scaffold Library. The use of a scaffold library to produce ligands with a known structure was validated by identifying peptides that bind to an anti-lipopolysaccharide (LPS) monoclonal antibody. A library was synthesized into a highly structured peptide that binds zinc in its three-dimensional conformation. PYKPECFSFSQK(X/S)(X/D)L(S/V)(X/K)HQ(X/R)THTG, presenting the randomized (X) residues to the target molecule on a defined backbone. Because of the peptides zinc binding, the conformation of the peptide mixtures and individual peptides could be confirmed by Co(II)-complex fluorescence analysis, circular dichroism measurements, and transfer NOE NMR. A positional scanning approach was employed, synthesizing five libraries of the format O1X2X3X4X5, X1O2X3X4X5, X1X2O3X4X5, X1X2X3O4X5, X1X2X3X4O5, ~26 pools each (17 proteinogenic, 9 nonproteinogenic amino acids), where the O positions were defined according to the pool, and X represented mixtures. Substituting hydrophobic residues at X1 caused a major change in conformation, and thus only 14 of the amino acids were used at this position. Screening with the anti-LPS antibody produced H1F2V3Q4H5 as the best binder, and the backbone conformation of these residues was confirmed by physical studies. The authors propose this as a first step toward synthesis of a peptidomimetic and represents an approach that can be generalized for any target molecule.

In another example of the use of a template library, FmocK-P-G-BocK-A-AlocK-P-G-BocK-A was synthesized, cleaved from the resin, and cyclized to serve as a backbone to which amino acids could be attached selectively to the orthogonally protected lysine side chains. A single amino acid was attached to the two Boc-protected lysines, while eight different amino acids were attached, in separate sets of reactions, to each of the other orthogonally protected lysine residue side chains. The limited library that resulted was screened against an anti-lysozyme antibody (HyHEC-5) which recognized a defined discontinuous epitope of lysozyme. Library screening identified a tyrosine that was not present in the native epitope. A similar concept was employed to display four amino acids from a Kemp's triacid backbone.
Multiple-Synthesis Libraries. Binding to streptavidin was used to validate a surface display array-based library method called Pilot. The effect of the surface on the binding properties of the compounds assayed while remaining attached to the solid phase on which they were synthesized is an issue. Cass et al. attempted to address the surface presentation issue by exploring the binding of $^{125}$I-streptavidin to the sequence YGHPQGG synthesized onto various surfaces. They derivatized polyethylene sheets with a functionalized linker (diamino-substituted polyethylene glycol, Jeffamine ED-600) to which they covalently bonded carboxymethyl dextran (Pharmacia T500) to produce a surface coating that was amino functionalized by coupling tBOC-NH-CH$_2$NH$_2$ followed by cleavage of the protecting tBOC. Arrays of compounds were synthesized onto these thin-film sheets in defined positions to create reusable assay plates.

The strategy used for array design is interesting. A standard mixture of 16 selected amino acids was used (W, amino acids Y,F,D,E,K,R,Nle,V,G, DaLa,A,S,H,Q,P,DNap) together with limited positions in which a mixture of two defined amino acids was incorporated. Thus, for instance, an 8 $\times$ 8 array hexapeptide library was generated by creating all possible two-amino acid combinations of the amino acids used in the W mixture in positions 1 and 2, and 3 with Y/F, D/E, K/R, Nle/V, G/DAla, A/S, H/Q, P/DNap by (Y/F, D/E, K/R, Nle/V, G/DAla, A/S, H/Q, P/DNap), while the W mixture was used in positions 3, 4, 5, and 6. Probing with streptavidin produced VY. Generating the subarray to define positions 3 and 4 (same format) produced VYGF and VYHP. Generating the final sub-array to define positions 5 and 6 produced VYGFRQ and VYHPQF. Speculating that these two might overlap the authors combined the two motifs to produce HPQVYGFRQ, which was a strong binder compared to the individual peptides.

The identification of DNA binding peptides from a library was demonstrated by Kramer et al. As described above, they synthesized a XXO$_2$O$_2$XX library on a cellulose membrane sheet in the form of pools in a 19 $\times$ 19, O$_1$ by O$_2$, array using the double coupling of 0.8 equimolar (less than saturating) mixtures of amino acids. The cellulose sheet was incubated with a $^{32}$P-labeled, double-stranded DNA 15-mer, and analyzed by a Phosphoimager (Molecular Dynamics, Sunnyvale, CA). This led to the identification of a KK, KR, RK, RR consensus for the O$_1$O$_2$ amino acid pair. Considering the crucial role of protein recognition of DNA sequences for transcription, and the current attempts to influence disease states through the selective control of transcription, these results demonstrate the potential for applying peptide combinatorial libraries to these targets.

Split-and-Mix, One-Compound–One-Bead, Postassay Identification Format Libraries. Lam et al. synthesized a pentapeptide library and assayed it directly for streptavidin binding to the compounds displayed on the beads, sequencing to identify the positive compounds, which were then resynthesized and tested to confirm that binding was competitive with biotin. The consensus motif...
identified was HPQ, with a lesser number of HPM motif compounds. Furthermore, it did not matter if HPQ was N-terminal (e.g., HPQFV), internal (e.g., LHPQF), or C-terminal (e.g., MYHPQ); of course, the C-terminal position in the library was internal to the linker. The HPM motif containing peptides was displayed either as an internal motif (e.g., MHPMA) or C-terminal motif.

As a follow-up, Lam and Lebl explored the specificity of streptavidin and avidin using a library that lacked histidine. A higher concentration of the streptavidin/alkaline phosphatase conjugate had to be used, but the result was the discovery of four motifs, WPAX or WXAX, RMDLY, and YMEXW, and two additional sequences, PPWPY and QYWQS. Representative compounds that were resynthesized demonstrated competitive binding with biotin, indicating they bound in the biotin binding pocket. This demonstrates the power of synthetic combinatorial libraries in discovering novel leads. Furthermore, when the apparently closely related avidin molecule was used as a target the authors found apparent overlap with streptavidin motif (HPYPX, HPFXX, and HPXPX motifs) as well as a different peptide specificity (HKXXX). Despite the apparent overlap, streptavidin did not bind to HPYP, and avidin did not bind to the LHPQF, indicating that the HPYP motif is specific for avidin and the HPQ motif is specific for streptavidin, though both bind in the respective biotin binding pockets.

Following this study additional L- and D-amino acid peptides were discovered. The results show that the majority of the L-amino acid-containing peptides that bind streptavidin contain the HPQ motif, while avidin binds D-amino acid containing peptides that do not have this motif, but instead exhibit a vynxx, ltxsx, or vqsxw motif, again demonstrating the binding pocket specificity differences that existed between the two biotin binding proteins.

Protein–protein and protein–DNA binding interactions regulate signal transduction, transcription, and translation. SH3- and SH2-mediated protein–protein interaction is an important component in signal transduction, serving to interface selective ligand/receptor interactions into common cellular signal transduction pathways. Specific inhibition of SH3-dependent binding is therefore an attractive therapeutic target. Studies were undertaken utilizing a hexapeptide library and a cyclic heptapeptide library assayed on-bead using the fluoresceinated SH3 domains of phosphatidylinositol 3-kinase (P13KSH3) and Src presence of at first all 32 peptides. The set was divided into two pools of 16 and the active pool was identified, then this pool was split in half, and so on. In the end one high-affinity ligand was identified, Ac-L-Lys-D-Ala-D-ALa. The authors demonstrated the utility of this approach, but also evaluated the pool size that could be employed, concluding that the method was useful for pool sizes of 100–1000 compounds. Many methods can identify activity from a pool of compounds, but capillary electrophoresis gives a direct determination of binding affinity and does not require labeling of compound or a specific biological activity.
Identification of Enzyme Substrates

Peptide substrates have proven to be good starting points for developing substrate-based inhibitors of enzymes, and synthetic combinatorial chemistry peptide approaches should produce leads that can then be used for rational drug design. The identification of enzyme substrates from libraries represents a method to improve screening assays and should prove useful in creating screening assays for putative orphan enzymes, which have no known substrates. Such a tool should be instrumental in characterizing and validating new therapeutic targets through identification for substrates for the classification of orphan enzymes, identification of putative substrates from protein sequence database searches based on the sequences identified from library screening, and identification of putative substrates for kinases involved in critical functions such as signal transduction.

Multiple Synthesis Libraries. A multiple-synthesis mixture strategy was used to synthesize a library of 19 Kemptide-related peptides (format L(R/X)RASL, R is the native Kemptide residue) and a library of 19 analogs based on the v-Src autophosphorylation site (RRLIEDA(E/X)WAARG, E is the native v-Src residue). The cleaved peptides were incubated with [γ-32P] ATP plus either the cAMP-dependent kinase or the v-Abl protein kinase, and the phosphorylated compounds were identified by phosphopeptide-selective liquid LCMS. When synthesis reached the point of randomization the resin was divided into four pools accounting for 16, 10.5, 31.5, and 42% of the total. Each pool was reacted with a different mixture of amino acids (I,V,T; Q,H; Y,D,E,K,R,P; F,G,A,L,M,S,H,W) so that the same proportion of reactive sites were occupied by equimolar amounts of each amino acid. The compounds were cleaved and the pools were recombined into one mixture of 19 compounds. Quality control was performed by MS, and the libraries were screened against their respective target kinases using [γ-32P] ATP. After removal of the enzyme by molecular weight centrifugal filtration (Microcon-10 unit), the peptide mixture was subjected to liquid chromatography. The recovered peaks were split for analysis of 32P incorporation by γ counting and specific MS detection of phosphopeptides.

Kemptide was the principle peptide phosphorylated by the cAMP-dependent kinase, but each analog was phosphorylated to some degree. The intensities of the negative ion (phosphorylated peptide, detecting the PO2− and PO3− ions) versus positive ion (peptide itself) permitted the authors to compute the percent conversion to phosphorylated peptide and rank them (R > H > G > Q = K > E > Y > H > A > P > T > T > S > (xle) > F > (xle) > W > V > M > A) according to suitability as substrates (MS cannot distinguish Ile and Leu, shown as xle). This ranking was consistent with that reported from a traditional study (R > H > K > A) but contained much more information. The study with v-Abl protein kinase identified the Leu or Ile analog as the major substrate, with the Val analog as a minor substrate.
Resynthesis and testing of the Leu and Ile analog peptides compared with the Src substrate Glu peptide demonstrated a preference for the Ile analog as substrate (Glu: $K_m = 1.56 \text{ mM}$, $V_{max} = 27.8$ nmol/min mg$^{-1}$; Leu: $K_m = 0.727 \text{ mM}$, $V_{max} = 20$ nmol/min mg$^{-1}$; Ile: $K_m = 0.67 \text{ mM}$, $V_{max} = 73$ nmol/min mg$^{-1}$). Thus libraries were shown to be effective means of defining substrates for both a receptor-dependent protein kinase (cSMP dependent kinase) and a receptor-independent protein kinase (v-Abl).

In a specialized variation of multiple synthesis, Songyang et al. synthesized a biased library MAXXXXXXXXAKKK, where X represents 15 proteinogenic amino acids (excluding C, W, Y, S, and T), and then cleaved this library into solution. The mixture was incubated with $p60^{src}$ protein tyrosine kinase and $[g^{-32}\text{P}]ATP$ and the phosphorylated peptides were isolated from the mixture by ferric chelation chromatography and identified by sequencing. The authors identified the EEIYGEFF motif.

**Split-and-Mix, One-Compound-One-Bead, Postassay Identification Format Libraries.** Wu et al. identified peptide substrates serin/threonin cAMP-dependent kinase, known to exhibit specificity for peptides containing an RR(X)S motif. A library of peptides displayed on the surface of beads were incubated with the enzyme in the presence $[g^{-32}\text{P}]ATP$. Substrate peptides were phosphorylated, all the beads were spread in agarose to immobilize them, and autoradiography was performed to locate the labeled beads. The $^{32}$P-labeled beads were recovered and identified. Substrates that identified by this procedure had higher affinity than the commercial synthetic cAMP-dependent kinase substrate, Kemptide (LRRASLG, $K_s = 1.33 \text{ mM}$). From approximately 500,000 compounds assayed, 55 positive compounds were selected from a pentapeptide library. Sequencing of two yielded the sequence RRYSV ($K_s = 0.67 \text{ mM}$). Sequencing three compounds out of 60 positive beads from a hepta-peptide library produced SQRRFST ($K_s = 0.83 \text{ mM}$), YRRTSLV ($K_s = 1.43 \text{ mM}$), and IIRRKSE ($K_s$ not determined). Two of these exhibited higher affinity for the enzyme than the commercial peptide.

The same approach was used subsequently to identify substrates for the $p60^{src}$ kinase. The sequence YIYGSFK was identified from the library. The phosphorylation of YIYGSFK and several peptides derived from cellular proteins by Src and several other kinases was studied. YIYGSF was found to be a highly specific Scr-family kinase (Scr, Lyn, Fyn) substrate, exhibiting a $K_m$ of 55 mM for Src, compared to a $K_m$ of 353 mM for a peptide derived from cdc2 (residues 6–20, KVEKIGEGTYGIVYK), a native substrate of Src, but not a good substrate for non-Src family protein kinases (EGFR, Fes) compared to the cdc2 peptide. Thus both a higher affinity and more specific substrate was identified.

The next step would be to search sequence data bases to identify whether this motif appears in cellular proteins that may be putative substrates for Src. Information of this type can be used to design substrate based inhibitors.
More recently, by screening a secondary library (XIXXXXX) an even more potent and specific peptide substrate, GIYWHHY, was identified.115

In another study the substrate specificity of an endoproteinase was explored using an acetylated library of peptides on beads.116 Two libraries were prepared. Ac-NleELRTQ(Pi)(P2)SHR-NH2, with P1 fixed as phenylalanine and P2 randomized in the first library, and P1 randomized with P2 fixed as serine in the second library. Incubation of the library beads with the 3C endoproteinase from hepatitis A virus resulted in removal of the acetyl group from those peptides that were cleaved by the proteinase. The absence of the acetyl group permitted direct identification of the substrate sequence by Edman degradation. Thus there was no selection step, only incubation, assessment of the extent of reaction by measuring free amino groups using fluorescamine, and then sequencing of the entire batch of beads. The native P1/P2 residues inferred from other studies are serine/phenylalanine, which demonstrated a $K_{cat}/K_{m}$ value of 0.004. The library results indicated a P2 preference for alanine, serine, and glycine, and a less selective P1 preference, but peptides containing a P1 arginine or proline were not cleaved. All the peptides that were cleaved were less active than the putative substrate domain, exhibiting $K_{cat}/K_{m}$ values of 0.1 to 1.0. Thus this study confirmed the putative cleavage specificity to be serine/phenylalanine for P1/P2.

The substrate specificity of human fibroblast collagenase and stromelysin was evaluated using an on-bead library approach in which a fluorescent reporter was cleaved into solution, resulting in solution fluorescence in wells containing positive substrate beads.117-119 Positive beads were recovered and the remaining material was microsequenced to determine the cleavage site. The first assumption was that the library should encompass a tripeptide window displayed in an alanine frame, AAXXXAA. To design the optimal method of covering this window an alanine scan was performed on the peptide G4P,L,A IL," cleaved by both collagenase and stromelysin. In terms of conventional nomenclature shown as subscripts to the amino acid residues, the P1 proline, P2 leucine, and P3 leucine, and to a lesser extent the P3 phenylalanine were important for either activity or selectivity. The authors concluded that PXXL was crucial for collagenase recognition as a substrate, and designed a library of three subforms, all synthesized on controlled pore glass (CPG):

\[
\text{Cop-A1A2A3X5B5A}\rightarrow\{\text{e-aminocaproyl}\},-\text{BAla-(3-aminopropyl)-CPG}
\]
\[
\text{Cop-A1A2X5A3B5A}\rightarrow\{\text{e-aminocaproyl}\},-\text{BAla-(3-aminopropyl)-CPG}
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Cop is a fluorophore (7-hydroxycoumarine-4-propanoyl) developed specifically for capping peptides synthesized on solid phase.112 A represents fixed alanine residues, X represents positions varied one at a time between 20 amino acids such that the identity of the residue was known from the position in the synthetic/assay array, and B represents a degenerate position containing 20 different amino acids. Thus each subformat library consisted of 400 pools in
which the identity of the X residues was known. The 20 proteinogenic amino acids were used, substituting S-methylcysteine for cysteine.

While a more powerful library design/identification strategy would have made it possible to simply synthesize and screen the complete tetrapeptide library, the results obtained with this approach were quite interesting. By defining cassettes of related compounds based on the shared identity of the N-terminal X residue, the relative importance of the C-terminal X residue could be compared. The cassettes defined as L, M, N, and P exhibited collagenase substrate activity. The L, M, and N cassettes exhibited an identical C-terminal X profile of U > L > M > I > Q > Y, while the P cassette had a different profile, N > U > A > D = M > P. Sequencing was used to identify the B residue identity and cleavage site. For instance, sequencing of the positive AAPANBA pool produced an abundance of and preference for B of U > L > I > M in the first round of sequencing, indicating that cleavage occurred N-terminal to these residues, and identifying AAPANUA as a good substrate for collagenase. Additional analysis indicated the ability to detect frame shifts in the point of cleavage. The authors also confirmed that there was a good correlation between cleavage of the peptides on the bead compared to cleavage of resynthesized peptides in solution. This approach should prove to be quite useful in the characterization of putative proteases identified from gene sequencing efforts and in the optimization of screening assays.

Another strategy used to identify protease substrates relied on an internally quenched fluorescent molecule where the fluorochrome (o-aminobenzamide, ABz) was separated from the quenching moiety (3-nitrotyrosine, Y(NO₂)) by the point of cleavage. By synthesizing the library on macroporous beads and employing on-bead screening, the beads that displayed substrate peptides developed fluorescence and could be readily identified and selected. The format of the library was Y(NO₂)X₁X₂PX₃X₄X₅K(ABz) bead, designed for the endopeptidase subtilisin Carlsberg. By inserting proline at the P-2 position the site of cleavage by this enzyme was directed between X₃ and X₅. By performing the assay at pH 8.5 and 5.2 the authors demonstrated that they could not only identify substrates, but they could optimize substrates for different reaction conditions. The substrates with the highest rates of cleavage on beads at pH 8.5 were Y(NO₂)LQPFNEK(ABz), Y(NO₂)FSPLQFK(ABz), and Y(NO₂)IEPFFEK(ABz), while those with the highest rates of cleavage at pH 5.2 were Y(NO₂)FQPLDVK(ABz), Y(NO₂)LYPLDVK(ABz), and Y(NO₂)ASPMGFK(ABz). However, the authors noted that while reflecting the rate of cleavage, this rank order did not hold in solution where accurate kinetic constants (Kₐ/Kₐ) could be determined. Thus, final optimization had to be performed using compounds assayed in solution, producing Y(NO₂)FQPLDEK(ABz) and Y(NO₂)FQPLAEK(ABz) as the most active structures under pH 8.5 conditions.

**Iterative Synthesis and Screening Library Methods.** Phosphorylation of library peptides was also used to identify substrates of cAMP- and cGMP-
Validation of Library Design Concepts. An early study in which enzyme substrates were identified was based on the synthesis of several mixtures of peptides. Each mixture contained compounds of identical structure except at one position, which was randomized (e.g., Ac-GNSXR, where X = F, G, A, Y, L, V).\textsuperscript{14} The mixtures were then exposed to an enzyme (angiotensin-I converting enzyme, atrial dipeptidyl carboxyhydrolase, bacterial dipeptidyl carboxyhydrolase, and meprin), and the products were analyzed by sequencing to determine which residues were important for binding and where the point of cleavage occurred. One cycle of sequencing revealed the preferred \( \textit{P}_i \) residues, two cycles the preferred \( \textit{P}_j \) residue, and so on. The authors made several new observations regarding substrate specificity, but the most interesting was perhaps the observation that meprin exhibited specificity for the Ala-Pro bond, while also cleaving Phe-Ala bonds. The authors speculate that bradykinin may be a physiological substrate of meprin.

Identification of Enzyme Inhibitors

Multiple Synthesis Libraries. Inhibitors of acetylcholinesterase activity were identified using a library of compounds synthesized and then spotted on a silica thin-layer chromatography plate and assayed.\textsuperscript{24} Fifty-two compounds were synthesized and spotted in an array. The substrate acetylthiocholine iodide plus 5,5-dithiobis(2-nitrobenzoic acid) and then enzyme were evenly distributed over the plate by spraying. Inhibitors were detected by the clear zones that appeared because of the lack of enzymatic reaction (which produced a yellow background). Cellulose paper could also have been used as the support, in which case synthesis could have been carried out directly on the support. Positive control and several other inhibitors were identified, validating the method.

Synthesis of a library based on a noncleavable peptide inhibitor on pins was used to map the active site of and optimize inhibitor selectivity for human heart chymase, a chymotrypsin-like protease that converts angiotensin I to angiotensin II.\textsuperscript{25} Two libraries were synthesized and the compounds were tested for inhibition of either chymotrypsin and chymase. One was based on use of 3-fluorobenzylpyruvamide (F-Phe-CO-) to bind in the \( \textit{P}_i \) pocket of the
enzyme. It consisted of (F-Phe-CO)-X,X,-RG-pin, where X,X,-residues were each varied among the 20 different proteinogenic amino acids to produce 400 different peptides used to determine the specificity for the P1 and P2 residues. This library identified the X,X,-residue pair ED as selective for chymase. Synthesis and testing of (F-Phe-CO)-ED-RG(OMe) resulted in the determination of a KI for chymase of 1 mM and for chymotrypsin of 100 mM. The other library was based on the use of an α-ketoamide (Phe-CO) in the P1 position, producing the library Z-Ile-X,X,-(Phe-CO)-GG-pin, where X,X,-residues were each varied among 18 of the different proteinogenic amino acids (Met and Cys were omitted) to produce 324 different peptides used to determine the specificity for the P1 and P2 residues.

The results were the identification of chymase P1,P2 preference for the amino acid pairs E2P2 and I3Q2. E2D2 was the most potent inhibitor, but it was not selective. The authors selected E2P2 for its acceptable potency and selectivity, synthesized the peptide Z-IEP-(Phe-CO2Me) and established a KI of 1 nM for chymase and 10 nM for chymotrypsin. Clearly, the potency of these two series were quite different, but selectivity of either 100- or 10-fold could be achieved. The authors then combined these series into one molecule, Z-IEP-(Phe-CO)-EDR-OMe, which exhibited a KI for chymase of 100 nM and a KI for chymotrypsin of 40 mM, a 400-fold selectivity for chymase with reasonable inhibitory potency. This demonstrates the power of rationally designed combinatorial peptide libraries to quickly optimize the selectivity of enzyme inhibitors.

Split-and-Mix, One-Compound–One-Bead, Postassay Identification Format Libraries. Meldal and Svendsen described a novel approach for identifying enzyme inhibitors. The authors synthesized an intramolecularly quenched (by Ac-Y(NO2)] fluorescent (Abz) substrate for subtilisin Carlsberg (Ac-Y(NO2)FQPLAVK(Abz)-PEGA, cleaved at the L-A bond) inside macroporous beads composed of polyethylene glycol cross-linked polyamide (PEGA) together with a library of heptapeptides (XXXxxXXVF, composed of the L (X) or D (x) stereoisomers). Enzyme inside the beads thus was in the presence of both a substrate and a potential inhibitor. Cleavage of the substrate present in each and every bead produced fluorescence. However, in those beads containing an inhibitory peptide the development of fluorescence was delayed due to the localized inhibitor. Novel inhibitors were identified, the most potent being AMMc(tert-butyl)MIVF (IC50 ~ 3 mM).

The identification of inhibitors of protein tyrosine phosphatase PTP1B (implicated in breast and ovarian cancer) was used to validate a novel tagging method employing a radio-frequency (rf) encodable microchip to encode the synthetic steps to which a compound carrier system was exposed in a split-and-mix, one-compound–one-carrier approach to library synthesis (See Chapter 14). A tripeptide library acetylated with p-carboxycinnamic acid (a non-specific inhibitor) was synthesized on solid-phase resin contained in capsules that also held an rf microchip emitting a unique binary code. The objective
was to confer specificity and additional binding affinity through the peptide interaction with enzyme residues surrounding the \( p \)-carboxycinnamic acid binding pocket. Each reaction pool was scanned and the identity of the microchips in that pool was recorded. The capsules from the various pools could then be mixed, split into the next set of reaction pools, and scanned again. Thus, at the end of the synthesis there was a record of which reactions each microchip-encoded capsule had been subjected to. In the actual example, a limited library of 125 compounds was synthesized, so the split-and-pooling strategy was also manually tracked to verify the ability of the rf tracking to accurately report the reaction history of each capsule.

Inhibitors of the glucosomol phosphoglycerate kinase (gPGK) from *Trypanosoma brucei* were identified from a pentapeptide library in an on-bead assay using fluorescently tagged or biotinylated gPGK and either manual selection of the fluorescent beads or selection onto steptavidin-coated magnetic beads. Inhibitors of this gPGK are potential antiparasitic agents, because of the key role this enzyme has in the glycolytic pathway of the form of *Trypanosoma brucei* found in the bloodstream, and because it is known that the *Trypanosome* is sensitive to compounds that interfere with glycolysis. Using this approach the authors found that NWMMF was a selective inhibitor of the parasite gPGK, \( IC_{50} \) of 80 mM, and was without effect on rabbit muscle gPGK at 500 mM. The next step toward a therapeutic agent is to optimize activity of this parasite-selective peptide and demonstrate anti-parasite activity.

Lowe and Quarrell started with a portion of sequence from a known 29-amino acid residue inhibitor of the serine proteases trypsin and chymotrypsin, CMT-1 (isolated from squash) to design an optimization library for leukocyte elastase. The portion of sequence they used as a template defined a reactive loop in this conformationally constrained inhibitor. The authors synthesized a library containing mutants at positions \( P_1 \) to \( P_4 \) (C-terminal) to the presumed reactive site occurring between Arg-5 and Ile-6 relative to the N-terminus. Once cleaved this inhibitor remains bound to the enzyme active site. Other investigators had shown that changing Arg-5 to Val-5 increased binding to human elastase. This led the authors to design two libraries, differing in the fifth residue, \( RVCP(V/A)XXXXC_KKDSDCLEVCLEHGYGCG \), and randomizing the \( P'_1 \), \( P'_2 \), \( P'_3 \), and \( P'_4 \) positions with the 19 proteinogenic amino acids. The residues that induced the tertiary conformation of the peptide through disulfide bond formation were all retained. Using this template library and fluorescently labeled human leukocyte elastase, they screened a therapeutically relevant serine protease in an on-bead assay using fluorescence microscopy to select positive beads. Disappointingly, the authors did not actually report on the success of screening or identify the peptides found using this library.

Inhibitors of thrombin are known to bind in both the active catalytic site as well as an exosite. An example where this was used to advantage is the inhibitor Hirulog, \( lPRPGGGG(NorLeu)GDFEEIPEY \) \( (K_i = 2 \text{ nM}) \).
log was designed on a rational basis from a known peptide that constitutes a pharmacophore that binds in the active site of the enzyme (fPRPG, \(K_i = 20 \text{ mM}\)), a linker, and a sequence derived from a large molecular weight inhibitor, Hirudin, comprising a pharmacophore that binds in an exosite (DFEEIPEEYL, \(K_i \sim 2 \text{ mM}\)). Pentapeptide libraries were assessed using on-bead screening to identify active site inhibitors. From a library of format DLDLD, where a set of D- or L-amino acid isomers were randomized at every other position, several novel compounds were identified. Among them was wFrPf, which exhibited a \(K_i\) for thrombin of 3.5 mM, and dYaEw (\(K_i = 6 \text{ mM}\)), compared to a \(K_i\) of 21 mM for the known pentapeptide inhibitor fPRPG (the \(n\)-terminal sequence of Hirulog, a thrombin inhibitor in clinical trials). From a library of the format LDLDL compounds such as TrFfP (\(K_i = 117 \text{ mM}\)), LiFrP (\(K_i = 20 \text{ mM}\)), and GfFfK (\(K_i = 23 \text{ mM}\)) were identified, while from a library of the format DDDDD compounds such as ffrfh (\(K_i = 23 \text{ mM}\)), ffrpt (\(K_i = 43 \text{ mM}\)), ffr(norleu)(norleu) (\(K_i = 53 \text{ mM}\)), and rlmrf (\(K_i = 83 \text{ mM}\)) were identified.

One benefit of combinatorial libraries in exploring binding domains around an initial starting point is to identify a novel series, as well as to improve activity of the original. With this in mind an on-bead library was designed with the format fPRPXXXX-(linker)-bead, randomizing 19 proteinogenic amino acids at each of the X positions. Screening this library identified SEL 1478, FPRPFGYRV-NH\(_2\), which exhibited a \(K_i\) of 25 nM. Thus, using a single optimization library, the affinity of the peptide lead was optimized from 20 to 25 nM and a novel binding domain was identified. Multiple synthesis was employed in a combined alanine scan and truncation study to rapidly explore the SAR of SEL 1478, with the result that the compound FPRPFG(X)R-NH\(_2\) was identified as necessary for maximal activity. Optimization using an expanded library of nonproteinogenic amino acids and peptide bond modifications would be the next logical step in optimization.

Factor Xa is another enzyme in the coagulation cascade. Drug discovery teams have searched for a specific low molecular weight, orally available inhibitor of this enzyme for the last 5–10 years without success. However, by screening an octapeptide library in an on-bead assay a novel lead was identified, YIRLAAFT-NH\(_2\) (SEL 1691, \(K_i = 15 \text{ mM}\)). Many additional analogs were identified from other peptide libraries, including the consensus motif YIRLA, structurally different from any known peptide serine protease inhibitors. Furthermore, the series was specific for factor Xa. This factor Xa lead was optimized in a combined combinatorial chemistry and traditional medicinal chemistry one-at-a-time synthetic approach.

The first effort was the multiple synthesis of truncation and alanine substitution analogs, resulting in the identification YIRLA, as the structure necessary for activity. Libraries which allowed multiple substitutions within this motif were designed, synthesized, and tested. This provided valuable SAR and, as the most active compound, SEL 2060 (\(K_i = 150 \text{ nM}\)). Traditional medicinal chemistry approaches based on SAR and molecular modeling ultimately pro-
duced a series of very active and specific inhibitors, among them the pentapeptide SEL 2711, Ac-p-amidonophenylalanylcyclohexylglycyl-3(methyl pyridinium)alanyl-L-P-NH2. This molecule exhibited a $K_i$ of 3 nM for factor Xa, was >3000-fold selective compared to thrombin and a variety of other enzymes, was not cleaved by any serum enzymes, and exhibited antithrombotic activity in animals after intravenous, subcutaneous, and oral administration.

Iterative Synthesis and Screening Library Methods. Inhibitors of the HIV-1 protease were identified from a tetrapeptide library. The peptides were screened free in solution in a microplate assay in which the cleavage of a fluorescent substrate by the HIV-1 protease was monitored. The library contained 22 amino acids at each position plus the aspartic acid protease transition-state analog Statine (Sta) at the second position. Twenty-two pools were synthesized to define the first position as Ac-(P/D)-XXX-NH2. Successive rounds of synthesis defined the rest of the peptide to be Ac-PI(Sta)D-Leu/D-Phe)-NH2, the aspartic acid having lost activity/specificity on the second round of synthesis and screening. Thus, of the 23 residues incorporated at the second position, only the Statine-containing peptides were active, the activity ($IC_{50}$) of the most active peptide (Ac-PI(Sta)\textit{p}-Leu)-NH2 was 1.4 mM. Subsequent substitution of the $\text{p}$-leucine with amino acids not in the original library yielded three leucine-substituted peptides (L, V) with 10-fold greater potency; substitution of isoleucine with valine produced a 2-fold enhancement of activity. Reassessment of the valine series indicated that tryptophan was more active than phenylalanine at the N-terminus, producing the tetrapeptide Ac-WV(Sta)\textit{p}-Leu)-NH2, $IC_{50}$ 200 nM. Further optimization from the initial lead produced a pentapeptide with a defined composition Ac-WV(Sta)VX (X not disclosed) with an $IC_{50}$ of 5 nM.

Eichler and Houghten identified inhibitors of trypsin from an iterative set of 10 hexapeptide libraries with the P\textsubscript{i} position defined as K or R, synthesized in an array on cotton and screened in solution (Ac-KXXXXX-NH2, Ac-XXKXXX-NH2, Ac-XXKXX-NH2, Ac-XXXKX-NH2, Ac-XXXXKO-NH2, Ac-ROXXX-NH2, Ac-XROXX-NH2, Ac-XXROXX-NH2, Ac-XXXROX-NH2, Ac-XXXXRO-NH2). This strategy, incorporating 19 of the proteinogenic amino acids, resulting in 190 pools containing 2,606,420 peptides, produced the sequence Ac-AKIYRP-NH2, $IC_{50}$ = 46 mM, which was slowly cleaved by enzyme, but was a better inhibitor than the most active hexapeptide sequence derived from naturally occurring trypsin inhibitors (Ac-TKIYNP-NH2, $IC_{50}$ = 102 mM). The authors went on to design a secondary library to optimize the activity of this novel lead.

Lead optimization was pursued using the iterative process described above of synthesizing 400 pools of composition Ac-OiO3XXXX-NH2, screening, and then synthesizing and screening subsequent iterative sets of 20 libraries each (Ac-Id1Id2O3XXX-NH2, Ac-Id2Id3O3XX-NH2, etc.) and proved unsuccessful in identifying any inhibitors of trypsin. Eichler and Houghten synthesized an iterative set of 10 hexapeptide libraries with the P\textsubscript{i} position defined
(biased) as K or R, as described above, and identified a novel hexamer sequence, Ac-AKIYRP-NH₂, IC₅₀ = 46 mM. The authors incorporated this into the design of a secondary dodecamer library to optimize the activity of this novel lead. Through another series of iterative synthesis/screening steps the most active dodecapeptide was identified. An example of one library is Ac-XXXAKIYRPPOXX-NH₂. Altogether 19 libraries were synthesized in which the randomized position and the AKIYRP motif was varied positionally throughout the template as the KO motif had been positionally varied. The final result was Ac-YYGAKIYRPDKM-NH₂, with an IC₅₀ of 10 mM.

Petithory et al. had demonstrated that endopeptidase substrates could be identified from peptide libraries. The identification of selective inhibitors of metalloproteases, the rat brain zinc endopeptidases 24-15 and 24-16, which metabolize neuropeptides, was achieved using a phosphinic peptide library of the format Z-(L,D)PheY(PO₂CH₂)₃-(L,D)(G/A)₃-X₃X₅, where X was substituted by 20 amino acids (See Chapter 9). Phosphinic peptides (with in vivo activity) act as mixed inhibitors of the two peptidases, with nanomolar IC₅₀, without affecting several other zinc peptidases, such as endopeptidase 24-11, angiotensin-converting enzyme, aminopeptidase M, leucine aminopeptidase, and carboxypeptidases A and B. However by varying the P₁ position between G and A, and randomizing the P₂ (X₁) and P₃ (X₃) residues the authors identified selective inhibitors from their optimization library.

A single-step deconvolution strategy was used to synthesize the peptides. X₃ was coupled to resin in 20 different reactions, the resin was mixed and split, and then X₂ was coupled in 20 different reactions. Each of these mixtures was split and then either Z-(L,D)PheY(PO₂CH₂)₃-(L,D)G-OH or Z-(L,D)PheY(PO₂CH₂)₃-(L,D)A-OH was coupled. This scheme produced 20 pools containing 40 Z-(L,D)PheY(PO₂CH₂)₃-G . . . (X₃) G/A-X₂ each, with G and X₂ defined, and 20 pools containing 80 Z-(L,D)PheY(PO₂CH₂)₃-A . . . (X₅) peptides each, with A and X₅ defined. The peptides were cleaved from the resin and assayed. The X₅ selectivity determined for the Z-(L,D)PheY(PO₂CH₂)₃-G-X₁X₅ sublibrary was R/K for 24-15 and P for 24-16. Twenty Z-(L,D)PheY(PO₂CH₂)₃-GRX₅ peptides were synthesized, purified by HPLC, and tested, identifying X₅ selectivity for 24-15 to be F/M (~400-fold selective, IC₅₀ = 2.7/0.35 nM, respectively). Pursuing the Z-(L,D)PheY(PO₂CH₂)₃-(L,D)A-X₁X₅ series the authors identified Pro, Nle as unselective, and Arg, Lys as 24-15-selective X₅ residues. The deconvolution synthesis of X₃ peptides a nearly 2000-fold selective inhibitor was identified, Z-(L,D)PheY(PO₂CH₂)₃-(L,D)A-K-M, exhibiting a 24-15 IC₅₀ of 0.12 nM and a 1300-fold selective inhibitor was identified, Z-(L,D)PheY(PO₂CH₂)₃-(L,D)A-R-M, exhibiting a 24-15 IC₅₀ of 0.07 nM. This is a striking achievement of both selectivity and potency from an optimization library strategy.

**Positional Scanning Format Library Methods.** Inhibitors of chymotrypsin were discovered using a cyclic peptide template library. A peptide backbone (KKKEG-resin) was synthesized using the tea-bag method of synthesis. Three
different positional libraries were synthesized, requiring three different backbone structures using orthogonally cleavable protecting groups on the lysine residues, depending on whether the residue being coupled was to be defined (Dde) or consisted of a mixture (Boc). The peptide backbone was cyclized between the N-terminal lysine and the glutamate residue before coupling of 19 proteinogenic and 10 carboxylic acids to the side chains of the three lysine residues to generate three positional libraries, cyclo[K(X)K(X)K(O)E]G-OH, cyclo[K(X)K(O)H]K(X)E]=G-OH, and cyclo[K(O)H]K(X)K(X)E]=G-OH. In these libraries O represents positions defined for each pool, X represents positions in which the mixture of side-chain acylating residues were attached, and the square brackets represent the cyclization between the N-terminal K and C-terminal E residues. A novel series of compounds was identified exhibiting side chains of piperonylic acid and 2-thiophenecarboxylic acid, with \(-50\) mM IC\(_{50}\) values (compared to 1.6 mM for chymostatin and 35 mM for Ac-\(\text{ggyyy}-\text{NH}_2\)).

**Validation of Library Design Concepts.** Using a more constrained and well-defined cyclic peptide scaffold, Domingo et al.\(^{127}\) investigated the P\(_1\) specificity for a series of different proteases. Their approach was based on the synthesis of an 11-mer peptide \(\alpha\)-chymotrypsin inhibitor\(^{165}\) mimicking the larger 6- to 9-kD Bowman–Birk inhibitors. The Bowman–Birk inhibitors present a nine-residue binding loop to proteases spanning the P\(_3\) to P\(_0\) binding domain, while the peptide mimic presents a seven-residue loop to the enzyme. The authors synthesized a library (SCTX,SIPPOQCY) wherein each member had an identical sequence, randomized only at the P\(_1\) position (X\(_1\)) by the 21 proteinogenic amino acids to generate a mixture of compounds. They screened this mixture for binding to porcine pancreatic elastase, subtilisin BPN\(^*\), trypsin, \(\alpha\)-chymotrypsin, and enzymatically inactive anhydro-\(\alpha\)-chymotrypsin and anhydro-trypsin. Positive peptides were selected by passing the mixture over affinity columns composed of immobilized enzyme, and then the retained peptides were sequenced to identify the cleavage site and P\(_1\) residues. In each case two sequences were obtained, one for the intact, bound peptide, and one for cleaved peptide. Trypsin exhibited a preference for K,R; \(\alpha\)-chymotrypsin for F,Y,M,L,Nle; anhydro-\(\alpha\)-chymotrypsin for F,Y; and elastase for L,M. \(K_i\) values ranged from 9000 to 10 nM.

**Identification of Receptor Antagonists**

**Multiple-Synthesis Libraries.** The method of synthesizing individual peptides on an array of pins and then either directly assaying for activity while the peptides are still bound to the pins or after their release into solution, lends itself to exploring structure activity relationships (SAR) of analogs based on the sequence of a lead molecule. The approach was applied to exploration of substance P analog binding to the NK\(_1\) receptor.\(^{96}\) A radiolabeled binding assay was employed using a rat brain synaptosome tissue preparation. All the
peptides synthesized had been reported previously, the purpose of the experiments described in the initial publication being to demonstrate that the crude material released from the pins could be used to establish SAR without further purification. The same group continued with the approach, applying it to study the SAR of substance P binding to the NK1 receptor through the simultaneous synthesis of 512 analogs to cover all possible D- and L-amino acid combinations of 9 of the 10 native residues of substance P (the C-terminal Met residue was not varied because it was directly incorporated into the support).

This exhaustive SAR produced new insights and a rich database. Most interesting was that single changes were not predictive of dual changes. This is a fundamental difference in traditional medicinal chemistry and combinatorial medicinal chemistry approaches. In the traditional approach the parent molecule is frequently divided into domains and each domain is varied while the rest of the molecule is kept constant. Later on the best modifications of each domain are combined and the molecule is further refined. In the combinatorial approach, multiple changes can be introduced, and optimized simultaneously.

Much of the effort in synthetic combinatorial chemistry is now focused either on the synthesis of nonpeptide libraries or on the use of rational approaches to improve the design and utility of chemical libraries such as the design of a dipeptide library based on physicochemical properties of select amino acids. After observing that the successful conversion of several peptides into nonpeptide peptidomimetics occurred through the discovery of millimolar dipeptides (angiotensin converting enzyme inhibitor CCK-B and CCK-A selective receptor antagonists), the authors considered that just using the 20 naturally occurring amino acids and four N-terminal protecting groups would generate 1600 possible peptides. They designed and synthesized an information-rich library of 256 N-protected dipeptides, encoding the greatest possible diversity of physicochemical information. The minimum analog peptide set was used, plus various N-terminal blocking groups (the des-amino derivatives of three amino acids, A, G, F), and the C-terminus was amidated, providing structures with at least three potential pharmacophores, which were described by the $z$ scales calculated using the physical properties of 55 amino acids. The set of 256 compounds was composed of dipeptides formed from eight amino acids (D, K, S, F, L, W, T, and V).

This library was used to identify NK3 selective antagonists, followed by more traditional optimization incorporating multiple synthesis where appropriate. The lead from the library was Boc(S)F(S)F-NH$_2$ ($IC_{50} = 1550$ nM), where the S indicates the chirality of the Phe residues. This compound was somewhat specific for the NK3 receptor, exhibiting an $IC_{50} > 10$ nM for NK1 and NK2 receptors. Of methylated analogs, only the C-terminal derivative retained activity, producing Boc(S)F(R,S)$\alpha$MeF-NH$_2$ ($IC_{50} = 1520$ nM). Exploration of the optimal chirality for the $\alpha$-MeF and attempts to functionalize the N- and C-terminal groups produced Boc(S)F(R)$\alpha$MeF-(CH$_2$)$_2$NHCONH$_2$ ($IC_{50} = 16$ nM), exhibiting >700-fold selectivity for NK3 over NK1 and NK2, and greater potency than the reference compound; SuccDFNMeFGLM-NH$_2$. 
Split-and-Mix, One-Compound–One-Bead, Postassay Identification Format Libraries. This method has been successfully applied to discover peptides that bind to the gpIIb/IIIa platelet integrin receptor. A cyclic pentapeptide library containing both D- and L-amino acids was synthesized and assayed by competition of soluble gpIIb/IIIa binding to fibrinogen-coated microplates. The peptide antagonists identified from the assay of only 50,000 compounds (a subportion of the library) were CRCDC and CROdC.

The C5a receptor has been the target of drug discovery efforts for many years. For nearly 5 years a traditional peptide effort was conducted at Abbott Laboratories to discover receptor antagonists using a rational approach focused on the C-terminus of the C5a ligand. A series of peptides containing nonproteinogenic amino acids were identified, but all were agonists, typified by the structure FKA(D-cha)(L-Cha)r-OH (binding IC_{50} = 70 nM). A subsequent effort carried out at another company did succeed in identifying an antagonist from this series after a similarly extensive effort. This target thus represented a relevant model to compare the advantages of a combinatorial peptide library to traditional peptide approaches.

A library was designed based on an analog of the Abbott agonist (FKP(D-cha)(L-Cha)r-OH) in which the parent residue was coupled to 20% of the resin at each step of the synthesis, while the remaining 80% was split among the other residues (proteinogenic and nonproteinogenic amino acids) used for randomization and the C-terminal arginine was not varied. A library of 1,200,000 compounds was created on a double releasable linker designed to release the peptides as a free C-terminal OH group. Screened to identify only those compounds with a binding IC_{50} better than 50 nM, 37 compounds were identified (e.g., FL(Tic)(D-cha)(L-Cha)r-OH, (NMe-F)LY(D-cha)(L-Cha)r-OH, FLY(D-cha)(L-Cha)r-OH), among them were close analogs to the antagonist described by Kawai et al. In this case a single library not only provided new SAR, but was successful in converting an agonist into an antagonist. The library approach identified the same lead in a matter of a few weeks of synthesis and screening, which had taken years by a traditional approach.

Iterative Synthesis and Screening Library Methods. Three publications have addressed the discovery and optimization of endothelin receptor antagonists using combinatorial chemistry approaches. Exploring SAR, all possible (64) L- and D-amino acid combinations were assessed for a potent ETR_{A} hexapeptide ligand (Ac-O(Orn)DIW-OH, IC_{50} = 230 nM), resulting in the identification of three analogs with millimolar activity, two containing 5 D-amino acid replacements, and with Ac-(D-orn)Diw-OH exhibiting a comparable IC_{50} as the parent of 250 nM. The technique used was the split-and-mix synthesis of a 30,752-member tripeptide library (containing 31 or 32 residues consisting of proteinogenic or nonproteinogenic amino acids at each position) designed around a known ET antagonist, FR-139317. Thirty-one pools (of mixtures of 992 compounds each) defined by the identity of the N-terminal residue, were screened. The authors then resynthesized and screened successive libraries to
define the remaining two residues. Not only was FR-139317 identified, but extensive SAR around this molecule was obtained. Deconvolution of mixtures from a $19 \times 19$ combinatorial array, based on a similar but more potent hexapeptide ETR receptor antagonist (Ac-(D-BHG)LDIW, $IC_{50} = 8$ nM) investigated by Spellmeyer et al.\textsuperscript{139} not only produced extensive SAR, but produced several analogs with comparable affinity as the parent and one analog with higher affinity (Ac-(D-BHG)QDVIW-OH, $IC_{50} = 1.7$ nM).\textsuperscript{140}

Thus, all three independent studies demonstrated the power of combinatorial chemistry optimization libraries to identify potent compounds and to provide extensive SAR.

**Positional Scanning Format Library Methods.** To discover antagonists of the binding of IL-6 to its soluble receptor (sIL-6R), Wallace et al.\textsuperscript{141} synthesized a positional scanning pentapeptide library. The ligand, IL-6, was bound to microtiter plates and then the library peptides were added with sIL-6R. Bound sIL-6R was detected with a specific anti-receptor antibody. No antagonists were discovered, so the investigators proceeded to synthesize the same library in a branched, polycyclic configuration using lysine residues as the branching points to produce an octameric library. The branched lysine core is shown in Figure 3.13. The library was synthesized onto the eight free amino groups, such that each branch was identical. Thus, a multimeric or polycyclic library was created in which each compound consisted of eight identical pentamers displayed off the branched lysine core. Again positional scanning was employed, but this time antagonists were identified, which produced a positional consensus $(E/D/Y)(F/I/E)(Y/L/E/F/W)(F/I/W/Y)(W/Y/F)$ from which a possible 360 compounds could be derived. To simplify follow-up the degree of inhibition caused by certain pools was used to limit the residue selection, reducing the number of compounds synthesized in the follow-up to 36 peptides.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure3.13.png}
\caption{Polydentate library: Use of the solid phase and a highly branched linker complex of lysine residues to present multiple copies of the same peptide to the target, enhancing weak binding interactions. Shown is a multimeric "octodentate" library of compounds.}
\end{figure}
At a concentration of 45 mM, 20 of these produced greater than 50% inhibition, and dose-response curves were obtained for the 18 most active. These results indicated a consensus sequence of (E/D)(F/I)(L/Y)(I/F)W, with an IC₅₀ of 75 nM for the six most active peptides (EFL1W, EFYIW, EI1FW, E1YFW, E1YIW, D1YFW). Finally, the authors synthesized these as well as negative controls as tetramers and dimers. The tetramers exhibited similar activity as the octamers, but the dimers were inactive.

Ligands that bind the T-cell receptor were defined in order to assess the structural requirements for recognition of the MHC class 1/antigen complexes. The mouse MHC class I amorphs H-2K and H-2L were chosen for study because both prefer octapeptides. The authors designed an octapeptide library, synthesizing eight libraries, each containing 19 pools with the position being defined occupied by 19 proteinogenic amino acids (OXXXXXXX, X0XXXXXX, XXOXXXXX, etc.). The ability to bind these peptides was assessed by the restoration of MHC expression. All the libraries contained peptides that bound and stabilized the two MHC class I molecules. In addition, the library pools sensitized target cells for cytolysis by T-lymphocytes, and the authors were able to define the specificity for several T-lymphocyte clones. In this functional assay the individual peptides identified were active in the pico- to femtomolar range. For instance, for one T-cell clone (4G3) that recognized the chicken ovalbumin-derived SIINFEKL epitope, they identified a novel epitope (DIKVGIEF), which was just as active. Even of greater interest was the identification of T-cell epitopes that induced cellular responses but that were inefficient in binding the MHC molecules. These latter molecules would appear to bypass the need for MHC interaction. This work demonstrates the potential of using synthetic combinatorial peptide libraries to understand the immune response and autoimmunity, and represents another approach to the development of synthetic vaccines.

**Identification of Receptor Agonists**

*Iterative Synthesis and Screening Library Methods.* A novel agonist of the m-opioid receptor, Ac-rfwink-NH₂, was identified from a hexapeptide library by testing the peptides in solution in a traditional radiolabel binding assay employing [³H][d-Ala²,MePhe⁵,Gly⁷-ol]enkephalin and rat brain tissue preparations. The selectivity of this all-d-amino acid-containing peptide was high (Kᵢ for m₁ = 16 nM and for m₂ = 41 nM), compared to the s receptor (Kᵢ > 1500 nM) and the kᵢ (Kᵢ > 2000 nM), kᵢ (Kᵢ > 5000 nM), and kᵢ (Kᵢ = 288 nM) receptor subtypes. Functional m-opioid receptor agonist activity (reversed by naloxone) was demonstrated on guinea pig ileum where Ac-rfwink-NH₂ exhibited an IC₅₀ of 433 nM, compared to leu-enkephalin, which exhibited IC₅₀ of 246 nM. In contrast, Ac-rfwink-NH₂ was functionally 100-fold less potent at the s receptor than leu-enkephalin, measured as the inhibition of electrically evoked contractions or mouse vas deferens tissue. Finally, intrace-rebroventricular injection of Ac-rfwink-NH₂ (3 nmol) showed analgesic prop-
properties in a nociception tail-flick model, exhibiting a long duration of action (120 min). Noteworthy is that the analgesic properties of this compound are more potent and longer lasting than those of morphine. The analgesic effect of Ac-rfwink-NH₂ was reversed by the irreversible m-selective antagonist β-funaltrexamine. Given intraperitoneal, Ac-rfwink-NH₂ also exhibited analgesic effects that were equipotent with morphine and reversed by intracerebroventricular naloxone. This observation suggests that the compound has bioavailability comparable to morphine. Due to its novel structure, bioavailability, potent activity, and good duration of effect, this compound or close analogs are being pursued as analgesics in an ongoing preclinical program.

L-Amino acid hexapeptides were also identified using a similar strategy. While very potent at the m receptor, (Ac-RFMWMT-NH₂ Kᵢ 0.8 nM, Ac-RFMWMK-NH₂ Kᵢ 0.4 nM, Ac-RFMWMR-NH₂ Kᵢ 0.5 nM), these compounds were less selective at the s receptor (Kᵢ of 0.9, 5.6, and 7.4 nM, respectively) and the k₁ receptor (Kᵢ of 0.6, 0.4, and 1.4 nM, respectively) than Ac-rfwink-NH₂. Subsequently, Dooley et al. identified several unrelated m-opioid agonists series, YPFGFX-NH₂ (X = R, IC₅₀ = 13 nM), WWPKHX-NH₂ (X = G, IC₅₀ = 9 nM), and Ac-RRWYXY-NH₂ (X = M, IC₅₀ = 33 nM), as well as antagonist, Ac-RWIGWR-NH₂ (IC₅₀ = 5 nM). Interestingly, the truncated sequence WWPK-NH₂ exhibited an IC₅₀ of 17 nM, WWPR-NH₂ exhibited an IC₅₀ of 10 nM, and both retained >200-fold selectivity for m- over s- and k- opioid receptors. These activities compare quite impressively with DAMGO (IC₅₀ = 3 nM; 253- and 844-fold selectivity over s and k, respectively) and YGGFL-OH (IC₅₀ = 5 nM; 0.7- and 257-fold selectivity over s and k, respectively). Relevant to the use of the iterative approach, the authors observed that at early stages in the iteration the YGG motif was more active, but because they pursued less active pools as well, they were able to define the YPFGFX-NH₂ and WWPKHX-NH₂ series, which were much more active and selective that the best YGGFMX-NH₂ molecules (IC₅₀ = 28 nM).

**Split-and-Mix, One-Compound-One-Bead, Postassay Identification Format Libraries.** A novel double-release assay was described for the identification of agonists and antagonists of G-protein coupled receptors. After library synthesis, dry beads attached by surface effect to polyethylene film were exposed to gaseous TFA in a controlled manner to cleave ~4% of compound from each, and then neutralized with gaseous ammonium hydroxide. The released compound remained trapped in the microporous beads. For the cellular receptor assay, tissue culture dishes of confluent frog (Xenopus laevis) melanocytes, transiently expressing the bombesin receptor introduced by electroporation or expressing either the frog α-MSH receptor or the transfected human α-MSH receptor, were treated with 1 nM melatonin. They were then covered with a thin layer of agarose containing melatonin and, in the case of the α-MSH receptor studies, α-MSH. The polyethylene sheet displaying the library beads was inverted over this.

With the beads now in (agarose) solution the released compounds were
free to diffuse out, contacting the underlying melanocytes. In this cellular system the pigmentation of the melanocytes is controlled through variations in the level of cAMP, which in turn is regulated by the G-coupled (bombesin or α-MSH) receptor. Agonists increase pigment darkening, whereas antagonists decrease pigmentation. In the bombesin studies, the plates with positive responses were frozen and the polyethylene film was removed (leaving the beads imbedded in the agarose). Image analysis was used to locate the positive beads, which were then collected by aspiration, washed, and subjected to a second round of release and screening.

The library synthesized for bombesin was derived from the sequence of bombesin, residues 8-14 (YAVGHLM-NH₂), and had the composition X₁X₂VGHLX₃-NH₂. X₁ and X₂ were randomized with 19 l-amino acids, and X₃ was randomized with L, M, P, W to produce a 1444-member library. In the first round of screening 5000 beads were assayed and 500 were selected and reassayed. From this second round of screening 52 beads were selected and reassayed to produce four positive compounds. The beads producing these compounds were recovered and sequenced to determine their identity. The activities of agonists identified ranged from ~100 nM (WFVGHLM and WAVGHLM) to 10 mM (AWVGHLM), compared to bombesin itself, which exhibited an EC₅₀ of ~0.1 nM.

This study demonstrated an elegant method for identifying receptor agonists. The α-MSH study was directed toward the identification of antagonists. α-MSH is a 13-mer, while the library synthesized for screening was a tripeptide library released as 48 pools of Ac-X₁X₂X₃-NH₂ and 48 pools of X₁X₂X₃-NH₂, in which X₁ was defined for each pool. Iterative synthesis and screening of subpools was used to define X₂ and X₃, though it would also have been possible to recover the beads corresponding to underlying pigmentation changes and determine the identity of the compounds from information on the beads had a coding method been employed as well as either a multiple release pooling strategy or more sparse seeding of beads over the cell layer. From these studies the authors identified numerous antagonists, among them the most active was wRL-NH₂ (Kd = 63 nM). The potential for this approach is that any G-coupled receptor transfected into the melanocytes will be automatically coupled to the pigmentation response through stimulus-dependent effects on cAMP levels. Thus, this system represents a universal assay for a wide variety of therapeutically relevant receptors that is capable of identifying both agonists and antagonists.

Expanding on this work the same group synthesized three libraries based on the α-MSH-[5-13] sequence: a 7-mer (minus X₁X₂), 8-mer (minus X₁), and 9-mer library, with the format X₁X₂(δL-F)(δL-R)(δL-W)X₆X₇P₈V₉-NH₂. The approach used was split and mix, one compound—one bead, but split into 8 subpools depending on the chirality of the residues 3, 4, and 5 (e.g., LLL, DLL, LDL, etc.). Thus, by knowing the pool and sequencing the individual active peptide the authors could identify the compound including the chirality. Because their assay enabled them to assess the activity of the
compounds from individual beads, as well as to identify both agonist and antagonist activity, they could assess individual responses within pools and not worry that compounds would interfere or cancel due to the simultaneous presence of an antagonist or agonist within the same pool. They identified an antagonist MP(D-F)(R)(D-W)FKPV-NH₂ (IC₅₀ = 11 nM), a partial agonist MP(D-F)(R)(D-W)WKPV-NH₂ (EC₅₀ = 690 nM), and a full agonist FH(D-F)(R)(L-W)QKPV-NH₂ (EC₅₀ = 0.4 nM); the agonist activity of α-MSH-[5-13] itself was EC₅₀ = 2 mM.

Identification of Functional Inhibitors

Iterative Synthesis and Screening Library Methods. Synthetic combinatorial peptide libraries have been screened to identify antimicrobial agents. In several reports of this application an iterative approach was used to identify L-amino acid-containing hexapeptides that exhibited antimicrobial activity, as well as D-amino acid- and nonproteinogenic amino acid-containing peptides with or without acetylation of the N-terminus or other postsynthetic modification. A liquid microbial growth assay was employed, and from these studies the peptide Ac-RRWWCR-NH₂ was identified as the most active among the L-amino acid-containing peptides, exhibiting a minimal inhibitory concentration (MIC) of 3.4 mg/mL against Staphylococcus aureus. From the libraries containing 56 different L, D, and nonproteinogenic amino acids at each position, four tetramers were identified: (aFmoc-elys)WfR-NH₂, (aFmoc-elys)WfL-NH₂, (aFmoc-elys)Wf(aAIB)-NH₂, and (aFmoc-elys)Wf-NH₂. The latter exhibited specific bacteriostatic activity for gram-positive bacteria.

Positional Scanning Format Library Methods. A hexapeptide positional scanning library was also used to identify peptides that inhibited the ability of melittin to lyse red blood cells. From a library containing 52,128,400 peptides, this study identified Ac-IVIFDC-NH₂, with an IC₅₀ of 11 mg/mL. This compound had no similarity to melittin and inhibited hemolysis in a competitive manner, which was overcome by raising the concentration of melittin. The authors suggested that the effect of the peptide is through a specific interaction, perhaps with melittin itself, rather than through a nonspecific membrane effect.

Split-and-Mix, One-Compound-One-Bead, Postassay Identification Format Libraries. A novel antimicrobial assay has been described based on pH-sensitive single release. The method was validated by use of analogs of the antimicrobial compound Ac-RRWWCR-NH₂, described originally by Houghten et al. Multiple synthesis of an alanine scan series and N- and C-terminal truncations of the peptide rrwwcr identified the importance of a C-terminal cre motif. There was only a marginal effect of deleting the N-terminal rr residues, and that the single replacements of the ww residues had the greatest effect on activity of any of the alanine replacements (the C-terminal...
DISCUSSION

Peptide combinatorial chemistry has confronted the medicinal chemist with the problem of how to carry out challenging reactions in a routinely predictable, automatable manner. A variety of solid supports have been explored and are generally available for the solid-phase synthesis of the various types of library formats. Similarly, much work has been devoted to overcoming the synthetic issues that are specific to library synthesis. Good methods exist to assure the equimolar incorporation of multiple subunits and to monitor reactions at the level of single beads, a necessity in cases of library synthesis, with the exception of multiple synthesis. Newer methods of structure determination have also been described, and with the advent of automation and deconvolution methods, this is often more an iterative or deductive process rather than an analytical challenge.

There are growing bodies of literature addressing cyclization of peptides, the scaffolds created to confer structural rigidity onto library members, and the functionalization of side chains; N- and C-termini, and the peptide backbone. One successful approach has been the use of an N-substituted glycine backbone, referred to commonly as peptoids (see Chapter 6).

The three most common synthetic and screening combinatorial approaches being employed are multiple synthesis, iterative/deconvolution synthesis/identification, and one-compound–one-bead synthesis and postassay identification. All three general approaches work, though in specific circumstances one approach may be the most appropriate. Multiple synthesis plays an important role in interfacing lead generation with lead optimization, but it can also serve a valuable lead generation function in situations where there is basis for good design, just as the potentially larger iterative or one-compound–one-bead libraries can play a crucial role in lead optimization and SAR generation. The advantage of a sophisticated one-compound–one-bead/encoding/sequencing
approach must be weighed against the investment that must be made in analytics for compound identification and in developing compatible chemical reactions and the expense of the linkers. Similarly, the elegance of a single synthesis must be compared with the need for iterative/recursive synthesis and screening, and the advantages of identifying an active compound versus elucidating the structure of a compound that should be active from the iterative algorithm. The synthesis and screening of mixtures is not just acceptable, but provides sound data and is in many instances crucial to success.

The research results that have been reviewed in this chapter demonstrate that combinatorial peptide approaches can identify peptide leads for many classes of targets that are of current therapeutic interest. For targets that have been extensively studied by traditional means, peptide combinatorial chemistry has frequently discovered novel leads and provided valuable new SAR. This in large part may be due to the ability to synthesize and screen so many compounds, and to incorporate many unnatural amino acids. It is also likely that such success is due to a difference between traditional and combinatorial approaches. In the former, medicinal chemists usually make one change at a time and assess the affects, while in the latter multiple changes are made at one time, permitting the chemist for the first time to efficiently explore the role of and synergistic intramolecular interactions.

There are fewer reports demonstrating the use of combinatorial peptide chemistry to optimize lead structures, but those that are reviewed here indicate the power of the approach. Identifying more leads is not the solution to all problems of drug discovery; it simply moves the bottleneck upward in the process to optimization. Recognition that combinatorial chemistry approaches can assist in lead optimization is fundamentally important to the combinatorial chemistry/drug discovery revolution. To realize the real benefits of combinatorial chemistry, the drug discovery team must think and apply the combinatorial paradigm through all phases of optimization, including animal testing.

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