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Integrin Ligands with α/β -Hybrid Peptide Structure: Design, Bioactivity, and Conformational Aspects

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* Tel: ++39-051-209-9450; Fax: +39-051-209-9456; E-mail: <u>luca.gentilucci@unibo.it;</u> web: http://www.ciam.unibo.it/gentilucci **Synopsis:** Integrins are cell surface receptors for proteins of the extracellular matrix and plasma-borne adhesive proteins. Their involvement in diverse pathologies prompted medicinal chemists to develop small-molecule antagonists, and very often such molecules are peptidomimetics designed on the basis of the short native ligand-integrin recognition motifs. This review deals with peptidomimetic integrin ligands composed of α - and β -amino acids. The roles exerted by the β -amino acid components are discussed in terms of biological activity, bioavailability, and selectivity. Special attention is paid to the synthetic accessibility and efficiency of conformationally constrained heterocyclic scaffolds incorporating α/β -amino acid span.

Keywords: β-amino acid; peptidomimetic; heterocycle; integrin; cancer

1. Introduction

Peptides fulfill a variety of biological and physiological functions, and for this reason natural and synthetic peptides have found a number of biomedical applications. Besides to the well-known bioactive peptides of natural origins, such as the anticancer, antibiotic, hormone, immunoactive, neuromodulator, and opioid peptides, new examples have been recently described for bio- and nanotechnological applications. Examples are peptide-derived vaccines, drug-delivery systems, radiolabeled probes, biomaterials for tissue engineering, self-assembling peptides, microchips, nanomaterials, biosensors, bioelectronic devices, and peptide-metal nanowires.¹

Unfortunately, many potential applications of native peptides remain unpractical, due to their limited stability *in vivo*, and to their poor ability to cross biological barriers (e.g. gut, blood-brain-barrier). Furthermore, peptides are rather flexible entities, which may result in lack of receptor selectivity. Therefore, much effort has been dedicated to the design of peptidic derivatives or analogues with improved stability, biological functions, and with the appropriate spatial orientation of the pharmacophoric groups. Among diverse strategies, the introduction of non-proteogenic amino acid residues is an effective and by now well-established method for the preparation of suitable peptidomimetics, i.e. compounds whose essential pharmacophore elements mimic natural peptides in 3D space, and which retain the ability to interact with the biological

targets.^{2,3} In this regard, within the many potential non-proteogenic amino acid residues to be used, the β amino acids⁴ have gained special attention.⁵ Indeed, there exists a significant number of naturally occurring β amino acids found in several natural products, for instance in fungal peptide antibiotics.^{3,6} β -Amino acids can be classified as β^3 , when the side chain is located adjacent to the β -amino group, or β^2 , when the side chain is adjacent to the carboxylic acid group (Figure 1). β^2 -Amino acids are generally less synthetically accessible relative to their β^3 -counterparts.⁶ From a conformational point of view, according to the convention of Banerjee and Balaram,⁷ the torsional degrees of freedom in a β -amino acid are defined as ϕ (N-C β), θ (C β -C α), and ψ (C α -C=O), respectively (Figure 1).

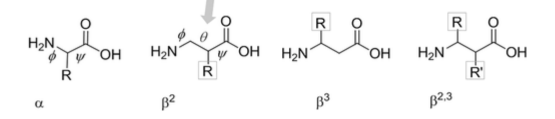


Figure 1 -Structures of α -, β 2-, β 3-, and β 2,3-amino acids (following Seebach's nomenclature⁶); the gray arrow shows the extra torsional angle θ (convention of Banerjee and Balaram⁷).

In spite of the extra C-C bond compared to α -amino acid residues, β -amino acids can afford well-defined secondary structures in β -peptides.⁸ Indeed, oligomers composed of all β -amino acids, or hybrid α/β -peptides^{9,10} have been extensively investigated for their tendency to adopt well-defined secondary structures,¹¹ in particular various types of helices.¹² Cyclic β -amino acids can further restrain peptide conformations as the C2-C3 bond becomes part of a ring.^{13,14} Folding oligomeric peptidomimetics are often referred to as "foldamers".^{15,16}

In this review, we focus on the use of β -amino acids and β -amino acid-derived scaffolds for the preparation of small-molecule integrin ligands with α/β -hybrid peptidic structure. Our aim is not to include all the available literature on the topic,^{5,6,10} but rather to discuss with the aid of specific examples the impact that β -amino acid residues have on biological activity, peptide stability, selectivity, and finally on three-dimensional (3D) structures. In particular, special attention is dedicated to the synthesis and use of heterocyclic scaffolds derived from β -amino acids. Very often, the preparation of scaffold-containing peptidomimetics requires multistep strategies, resulting in very low yields. Besides, many synthetic methodologies are not flexible and cannot provide structural and functional diversity in the target integrin ligands.

Furthermore, the conformational analysis of the β -amino acid peptidomimetics of interest appears to be fundamental for the comprehension of the chemistry and pharmacology of small-molecule integrin ligands. As a consequence, a central goal in this review is to show how the development of synthetic protocols for the preparation of conformationally constrained peptidic derivatives that maintain the required 3D disposition of the essential pharmacophores is currently of utmost relevance.

2. Integrins and their ligands

Integrins are heterodimeric, non-covalently bound proteins expressed on cell surfaces that mediate cell-cell and cell-matrix adhesion. These receptors constitute a family of 24 combinations of diverse α - and β -chains. Integrins regulate several aspects of cell behavior, such as cell proliferation, death, differentiation, and migration, 17,18 so they may represent privileged targets for therapeutic interventions. 19,20 For instance, $\alpha\nu\beta3$ integrin is responsible for cell adhesion to proteins of the extracellular matrix (ECM), a process involved in the construction of new blood vessels (angiogenesis). By contrast, the aIIbb3 (or GPIIb/IIIa) integrin plays a fundamental role in the aggregation of platelets mediated by fibrinogen. On the other hand, integrin $\alpha 4\beta 7$ is critical in lymphocyte adhesion to the gut. Integrin $\alpha 4\beta 1$ (VLA-4, very late activating antigen-4) is involved in cancer, diabetes and inflammatory diseases such as rheumatoid arthritis. It is responsible for the recruitment of leukocytes into inflamed tissues, and in the migration of activated lymphocytes during the immune response. In general, each of the 24 integrins has a specific function; however, a certain functional redundancy exists among integrins in some processes. For instance, the integrins $\alpha 2\beta 1$, $\alpha 3\beta 1$, $\alpha 9\beta 1$, $\alpha \nu \beta 5$, $\alpha v\beta 6$, $\alpha 5\beta 1$, are all up-regulated during wound healing. Beside, while some extracellular ligands are recognized by a specific integrin, other ligands show promiscuous interactions with diverse integrins; an example is represented by collagen, which can bind all integrins $\alpha 1\beta 1$, $\alpha 2\beta 1$, $\alpha 10\beta 1$, and $\alpha 11\beta 1$, albeit with different affinity.

The involvement of integrins in important pathologic processes inducted many research groups to block their functioning by means of monoclonal antibodies,¹⁹ or small-molecule antagonists of peptidic^{20,21} and non-peptidic structure.^{22,23,24} More recently, the new possibilities offered by nanotechnology led to the design of molecular devices functionalized with integrin ligands, to provide nanotools for integrin-mediated drug delivery,^{25,26} imaging,²⁷ or for theranostic applications.^{28,29,30}

The aim to develop specific integrin antagonists with appropriate pharmaceutical properties has been challenging. Many of the integrin superfamily members, including α IIb β 3, $\alpha\nu\beta$ 3, $\alpha5\beta$ 1, $\alpha\nu\beta$ 1, recognize the Asp-Gly-Arg (RGD) tripeptidic sequence (Figure 2A), an epitope shared by several proteins of the ECM including vitronectin (VN), fibronectin (FN), fibrinogen (FB), osteopontin, and bone sialoprotein.¹⁹ Other integrins interact with their ligands by adhesion to somewhat different recognition motifs. For instance, both α 4 integrins (α 4 β 1 and α 4 β 7) recognize a Leu-Asp-Val-Pro (LDVP) motif (Figure 2A) in the spliced CS-1 region in plasma FN, and a Leu-Asp-Thr-Ser (LDTS) sequence in the mucosal addressin cell adhesion molecule-1 (MAdCAM-1).^{19,31}

Recently, the use of antagonist molecules capable to target two or more integrins (in particular the RGDbinding ones) has been proposed.³² Indeed, it has been observed that cancer cells can modify integrin expression after drug treatment, and this might lead to the development of resistance to a single antagonist and, paradoxically, to promotion of tumor growth. For instance, the FN-binding integrins $\alpha\nu\beta3$ and $\alpha5\beta1$ are coexpressed on tumors and metastases, and each integrin regulates the functions of the other;^{19,33,34} the inhibition of integrin $\alpha\nu\beta3$ can increase cancer invasiveness in the presence of high levels of FN, due to increased recycle of integrin $\alpha5\beta1$, while inhibition of $\beta1$ integrins can promote tumor progression and metastasis by increasing the expression of $\beta3$.^{35,36}

In any case, selective integrin ligands remain fundamental in order to explore on a rational basis the 3D disposition of the pharmacophoric groups (e.g. the guanidino and the carboxylate groups of the RGD sequence). This systematic approach was termed "spatial screening" by Kessler et al.³⁷ However, because of the close similarities between the structures of the different members of the integrin family, and between their respective ECM ligands, the design of selective peptide therapeutic agents represented a hard task.³⁸ Illustrative examples of receptor-specific ligands are shown in Figure 2. The cyclic hexapeptide **1** c[Arg-Gly-Asp-D-Phe-

Val-Gly], or c[RGDfVG], were the lower-case amino acid abbreviation encodes a D-configured residue, is an efficient inhibitor of FB binding to the integrin α IIb β 3, IC₅₀ = 5.0×10⁻⁸ M, involved in platelet aggregation;³⁹ on the other hand, the cyclic pentapeptide **2** c[Arg-Gly-Asp-D-Phe-Val], or c[RGDfV], prevented binding of VN to the integrin α v β 3, IC₅₀ = 4.9×10⁻⁸ M, playing a role in tumor cell adhesion, angiogenesis, and osteoporosis. The different receptor specificity of these antiadhesive compounds was correlated to the different 3D disposition of their pharmacophores. Conformational analyses showed that the RGD motif occupies the positions *i*+1 to *i*+3 of a β-turn in the α IIb β 3 selective hexapeptide **1** (Figure 2B), while it resides in the positions *i* to *i*+2 of a regular γ -turn in the α v β 3 selective pentapeptide **2** (Figure 2C).³⁷ However, although it is highly selective for α v β 3 vs α IIb β 3, **2** is not selective over other RGD-binding α v integrins.

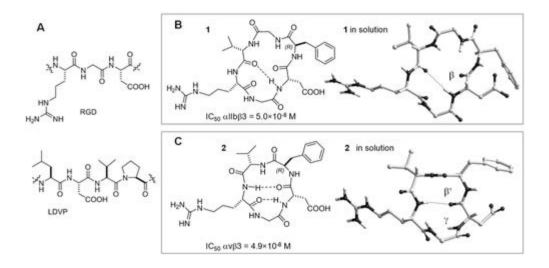


Figure 2 (A) The RGD and LDVP minimal binding motifs. (B) Structure and extended conformation in solution of **1**. (C) Structure and kinked conformation in solution of **2**. IC_{50} values refer to isolated integrins. The 3D structures were determined by two-dimensional NMR in combination with restrained and free molecular dynamics simulations.³⁷

Kessler and coworkers also investigated the biological activity of retro (DGR), inverso (rGd), and retroinverso (dGr) analogs (plus their stereoisomers) of the selective and superactive $\alpha\nu\beta3$ integrin inhibitor c[RGDfV] **2**.⁴⁰ Among the amino acid residues present in pentapeptide **2**, valine was found to be less important for binding $\alpha\nu\beta3$ and $\alpha\nu\beta5$ integrins.³⁷ For this reason, peptide c[RGDfK] become very popular since the lysine residue allowed to link the peptide over a length-optimized spacer and eventually to fluorescent tags, nanoparticles, or nanomaterials.^{25,28,41} Subsequently, each backbone amide segment in c[RGDfV] **2** was successively *N*-methylated to provide the monomethyl cyclopentapeptides. The effect of this derivatization on the activity was investigated, and revealed c[RGDf-*N*(Me)V] **3**, or cilengitide, as a highly active antagonist of the pro-angiogenic integrins $\alpha\nu\beta3$, $\alpha\nu\beta5$, and $\alpha5\beta1$, with IC₅₀ values of 0.65, 11.7, and 1.3×10^{-8} M, respectively (and a IC₅₀ of 8.1×10^{-7} M for the integrin α IIb $\beta3$), among the most potent ever developed,⁴² subsequently investigated in clinical trials as an anticancer therapeutic.^{43,44} In aqueous environment, **3** exists in a conformational equilibrium between two inverse γ -turns at Arg and Asp, and a γ -turn at Gly. The kinked backbone was identified as the most salient feature that differentiates $\alpha\nu\beta3$ and α IIb $\beta3$ antagonists.

The structural basis for the binding of $\alpha\nu\beta3$ to c[RGDf-*N*(Me)V] (**3**) was disclosed by the analysis of the Xray crystallographic structure of the extracellular portion of integrin $\alpha\nu\beta3$ in complex with **3**. The structure of the complex showed that the ligand is located at the interface between the $\alpha\nu$ and $\beta3$ subunits, resulting in extensive contact (Figure 3).⁴⁵ The $\alpha\nu\beta3$ integrin-c[RGDf-*N*(Me)V]-Mn⁺² structure (PDB ID: 1L5G) revealed that the peptide is inserted into a crevice between the propeller and β A domains on the integrin head. The five C α atoms of the cyclic peptide form a slightly distorted pentagon, and the side chains of Arg and Asp point to opposite directions. Strong ionic interactions bind the guanidinium group of Arg to Asp²¹⁸ at the bottom of the groove, and to Asp¹⁵⁰ at the backside. One of the Asp carboxylate oxygens is in close contact with the metalion-dependent adhesion site (MIDAS) in β A. The other carboxyl oxygen of Asp of the ligand is involved in hydrogen bonds with the amide protons of Tyr¹²² and Asn²¹⁵. Compared to the 3D model deduced by conformational analysis conducted in-solution for **2** and correlated cyclopeptides,^{37,38,40} the solid model of **3** shows some differences, being characterized by two inverse γ -turns centered on Asp and Arg, and a less 'kinked' RGD arrangement.

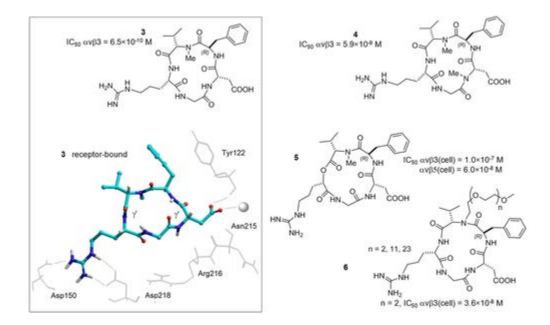


Figure 3 Structure c[RGDf-*N*(Me)V], cilengitide **3**; in the box: sketch of the main interactions of **3** with the $\alpha\nu\beta$ 3 binding site, as determined from the crystal structure of the complex (PDB ID: 1L5G); the ball represents a Mn2+ ion.45 Di-*N*-methylated **4**, depsi-analog of cilengitide **5**, and *N*-oligoethylene glycol substituted analogue **6**. IC50 values refer to isolated integrins or to integrin-expressing cells (cell).

The availability of the ligand-receptor X-ray crystallographic structure (see a sketch in Figure 3) gave the opportunity to analyze plausible candidate ligands by massive *in-silico* screenings, and to conduct highly detailed molecular docking analyses.⁴⁶ As for the other integrins, models of $\alpha 5\beta 1$ integrin were initially developed by homology on the basis of the $\alpha\nu\beta 3$ integrin-**3** complex structure,^{47,48} or by binding pocket mapping,⁴⁹ and were subsequently utilized to perform molecular docking simulations to rationalize the structural requisites of ligand-receptor binding. In a similar way, homology models^{50,51} or models derived from a combination of theoretical methods^{52,53} were utilized to design potential ligands of integrin $\alpha 4\beta 1$. Currently, high-resolution structural data are available also for the ectodomains of the integrin $\alpha 11b\beta 3^{54,55}$ and of the integrin $\alpha 5\beta 1^{56}$ in complexes with their respective RGD ligands. The "spatial screening" approach by Kessler and co-workers remains a fundamental step for the development of integrin ligand, but the conclusions derived from 3D conformational analysis not always agree with structural information provided by X-ray crystal structures of the integrins in complex with their ligands.

Since highly active and selective peptides are very useful tools to investigate the specific role of the integrin $\alpha v\beta 3$ in angiogenesis and cancer, Kessler et al. proposed di-*N*-methylated analogues of **3**. The di-methylation of the backbone consented to fine-tune the biological activity of the cyclic RGD sequences by stabilizing

 $\alpha\nu\beta3$ -specific bioactive conformations. In particular, the cyclopeptide c[RG-*N*(Me)D-f-*N*(Me)V] **4** (Figure 3) retained a nanomolar $\alpha\nu\beta3$ -integrin binding affinity, IC₅₀ = 5.9×10^{-9} M, but highly reduced activity for integrins $\alpha\nu\beta5$ and $\alpha5\beta1$, with IC₅₀ values > 3×10^{-6} M, and 2.7×10^{-7} M, respectively, and a IC₅₀ > 1×10^{-6} M for the integrin α IIb $\beta3$.⁵⁷

The structure of the prototypic ligand cilengitide **3** was the object of numerous modifications. For instance, Albericio et al. synthesized all five depsi-analogues of cilengitide by introduction of α -hydroxy acid residues in place of the normal amino acids. The analogue **5** (Figure 3) with ester link between *N*-Val and Arg showed increased potency with respect to the parent compound, being a potent small-molecule dual $\alpha\nu\beta3$ and $\alpha\nu\beta5$ antagonist in cell adhesion assays, IC₅₀ $\alpha\nu\beta3$ /FN = 1.0×10^{-7} M, IC₅₀ $\alpha\nu\beta5$ /VN 6.0×10^{-8} M, respectively, compared to IC₅₀ = 3.2×10^{-7} M and 2.0×10^{-7} M for cilengitide under the same conditions.⁵⁸ As another example, the *N*-Me group was replaced by *N*-oligoethylene glycol (N-OEG) of increasing length to provide the more lipophilic bioactive analogues **6** (Figure 3).⁵⁹

3. Integrin ligands with β -amino acid residues external to the recognition motifs

The rational design of integrin-selective peptidomimetic antagonists requires the precise prediction of the 3D structure. Herein we review the targeted incorporation of β -amino acids positioned externally to the recognition motif (e.g. RGD), and the consequent effects they exert on the geometry of peptide backbone. It has been shown that the β -amino acidic building blocks tend to adopt preferential conformations, and act as inducers of bioactive secondary structures. These findings can be utilized for designing peptidomimetic antagonists, since the specific 3D display of the pharmacophoric groups can be pre-defined by incorporation of tailor-made β -amino acids at a certain positions of the sequence.

3.1. Ligands with acyclic external β -amino acids or their mimetics

Sewald and coworkers have questioned whether cyclopeptides composed of both α - and β -amino acids may display distinct conformational preferences. In order to solve this question, the conformational impact of

the incorporation of a β -amino acid in model cyclopeptides was investigated by a systematic permutational replacement of one α -amino acid by the corresponding β^3 -analogue.⁶⁰ The cyclic pentapeptide c[Arg-Gly-Asp-D- β -Phe-Val] **7** (following Seebach's nomenclature,⁶ β -Phe in **7** corresponds to β^3 -Phenylglycine, β^3 -Phg) showed an high affinity for isolated $\alpha\nu\beta3$ integrins, IC₅₀ = 2.5×10⁻⁹ M, but in addition inhibited α IIb $\beta3$ integrin-mediated thrombocyte aggregation with an IC₅₀ value of 7×10^{-6} M, comparable to that of the reference cyclohexapeptide **1**. It can be appreciated that the β -amino acid residue is located at the central position of a tight inverse turn closely related to a γ -turn (Figure 4), namely a geometry whose dihedral angles can be correlated to a γ -turn conformation, the only difference being the insertion of a CH₂ into the peptide backbone (*pseudo* γ -turn). The authors observed that the superposition of the in solution-conformations of **1** and **7** reveal a satisfactory overlay of the peptide backbones and of the C α -C β vectors, accounting for the similar antagonist activities of the two peptides. On the other hand, pentapeptide c[Arg-Gly-Asp-D-Phe- β -Leu] **8** (β -Leu corresponds to β^3 -Val using Seebach's nomenclature⁶) presents a β II'-turn centered on Gly-Asp, with β -leucine placed in the central position of a *pseudo* γ -turn, suggesting that a β -amino acid may impose a stronger conformational control respect to a D-amino acid. This peptide showed improved affinity for isolated $\alpha\nu\beta3$ integrins, IC₅₀ = 9.6×10⁻¹⁰ M, and reduced efficacy against thrombocyte aggregation, IC₅₀ = 3.2×10⁻⁵ M.

Subsequently, Sewald et al. extended their studies to cyclotetrapeptides (CTPs) containing a β -amino acid residue, by introducing in c[RGDfV] **2** the β -amino acid D- β -Phe in place of the dipeptide D-Phe-Val. The cyclotetrapeptide **9** so obtained showed nanomolar affinity for the isolated integrin $\alpha \nu \beta 3$, IC₅₀ = 6.3×10^{-8} M, and a remarkable selectivity over α IIb $\beta 3$, IC₅₀ > 3×10^{-4} M. The β -amino acid residue is found in the central position of the pseudo γ -turn, thus inducing formation of the required γ -turn centered on Gly of the RGD motif (Figure 4). The conformation of ligands **2** and **9** seems indeed comparable and adequate for proper interaction with integrin $\alpha \nu \beta 3$ within the pharmacophoric regions.

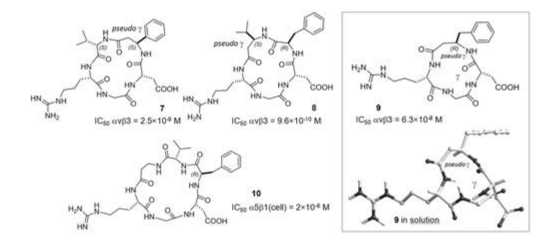


Figure 4 Cyclopentapeptides 7 and 8. In the box: the cyclotetrapeptide $\alpha\nu\beta3$ antagonist 9, rationally designed by incorporation of a β -amino acid as a pseudo γ -turn inducer; the RGD sequence forms a complementary γ -turn; 3D structure of 9 determined by 2D NMR, distance geometry, and restrained molecular dynamics simulations.60 Cyclohexapeptide $\alpha5\beta1$ antagonist 10. IC50 values refer to isolated integrins or to integrin-expressing cells (cell).

These preliminary results support the initial hypothesis that β -amino acids can act as inducers of *pseudo* γ -turns, allowing for the control of the peptide's backbone conformation, and consequently of the spatial orientations of the pharmacophoric groups of the RGD sequence.

The strategy was extended to the preparation of cyclic RGD peptides specifically designed for their affinity towards the integrin α 5 β 1. In particular, Sewald and co. synthesized a library of peptide sequences containing β -amino acids as inducers of the desired conformations. This search led again to the cyclopentapeptide **8**, and to the new cyclohexapeptide c[Arg-Gly-Asp-D-Phe-Val- β -Ala] **10** (Figure 4), low-micromolar inhibitors of the attachment of the α 5 β 1 integrin-expressing K562 cells to FN, with IC₅₀ values of 3×10^{-6} M and 2×10^{-6} M, respectively, compared to IC₅₀ = 2.8×10^{-4} M for the reference compound c-[Arg-Gly-Asp-D-Phe-Val] **2**.⁶¹ From the conformational analysis of the structure in solution of the peptide, it was concluded that a short average distance between the C β of Arg and Asp is required for binding to α 5 β 1.

Generally, cyclopenta- and cyclohexapeptides are easy to prepare, but their large rings imply high backbone flexibility. On the other hand, the smaller CTPs can be considered the minimal turn-mimics among the cyclopeptides. Unfortunately, CTPs composed of all α -amino acids are difficult to obtain, and frequently show mixtures of *cis/trans* isomers of the peptide bonds. Nevertheless, the incorporation of a distinct β^3 - or β^2 -amino acid residue in the sequence of the CTP can improve the yield of cyclization, and can stabilize the *all-trans* peptide bond conformation.^{60,62,63} In this context, partially retro cyclotetrapeptides (PR-CTPs) were proposed by Gentilucci et al.⁶⁴ as easily available CTP mimetics characterized by alternative secondary structures with respect to common CTPs (Figure 5). The cyclic structure and the presence of peptide bond surrogates was expected to improve enzymatic stability and molecular lipophilicity. The combination of *in vivo* stability and ability to cross biological compartments can strongly increase the bioavailability of drug leads based on these scaffolds. The stereoisomeric 13-membered PR-CTPs c[β -Xaa- ψ (NHCO)Ala- ψ (NHCO)Gly-Phe], containing a 1,2-diamine fragment as a β^2 -homo amino acid equivalent, were analyzed as models (Figure 5). The notation ψ (NHCO) has been proposed to indicate the occurrence of a retro peptide bond between two consecutive amino acids, i.e. -NH-C(O)- instead of the normal -C(O)-NH-.^{65,66}This implies that in cyclic partially retro peptide the first amino acid of the modified sequence is converted into a diamine and the last one is converted into a diacid (Figure 5).

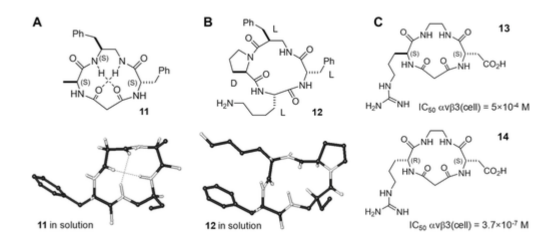


Figure 5 (A) Structure and backbone conformation of $c[\beta$ -Phe- ψ (NHCO)Ala- ψ (NHCO)Gly-Phe] **11**.64 (B) Structure and conformation of $c[(S)-\beta 2$ -Phe-D-Pro-Lys-Phe] **12**.63 Conformations determined by 2D NMR and molecular dynamics; some side chains have been omitted for clarity. (C) Structures of PR-CTP RGD mimetics **13**, **14**. IC50 values refer to integrin-expressing cells (cell).

Conformational analysis of the PR-CTPs models containing an unsubstituted diamine revealed certain backbone flexibility, especially in the diamine region, whereas PR-CTPs that incorporated a substituted diamine segment (e.g. **8**, Figure 5A) were much more rigid. In order to check if the PR-CTPs might represent novel 3D scaffolds respect to the 13-membered CTPs already reported in the literature, the conformations of $c[\beta-Phe-\psi(NHCO)Ala-\psi(NHCO)Gly-Phe]$ **11** (Figure 5A), and $c[(S)-\beta^2-Phe-D-Pro-Lys-Phe]$ **12**, reported by Arvidsson et al.⁶³ (Figure 5B) were compared.⁶⁷ These peptides share the same stereochemistry pattern: the (*S*)-diamine portion of **11** replaces the (*S*)- β^2 -Phe residue in **12**, L-configured Ala in the PR-CTP **11** corresponds to the D-configured Pro in the normal peptide, and the malonyl moiety ψ (NHCO)Gly adopts a conformation which is consistent to a D-amino acid residue. Nevertheless, the comparison of the two structures confirms that the two scaffolds are topologically non-equivalent (Figure 5), owing to the different hydrogen bonds, the position of the β -Phe side chain, below the macrocycle plane in **12** and above the plane in **11**, and because of the *cis* β -Phe-D-Pro ω bond in **12**.

The PR-CTPs were utilized for preparing the cRGD analogues c[β -Ala- ψ (NHCO)Asp- ψ (NHCO)Gly-Arg] (**13**) and c[β -Ala- ψ (NHCO)Asp- ψ (NHCO)Gly-D-Arg] (**14**) (Figure 5C), which were assayed as inhibitors of the adhesion of the native ligand FN to SK-MEL-24 (human malignant melanoma) cells, expressing the integrin $\alpha\nu\beta$ 3. The IC₅₀ values of **13** and **14** were 5×10⁻⁴ M and 3.7×10⁻⁷ M, respectively, clearly indicating that the different PR-CTP RGD possess distinct biological activities.⁶⁴

Moving from these preliminary results, the stereoisomers of the PR-CTP sequence $c[\beta$ -Phe- ψ (NHCO)Ala- ψ (NHCO)Gly-Phe] were analyzed as topology models having the three side chains directed towards precise directions. The topographic features of the eight PR-CTP models can be rendered using the Dunitz-Waser simplification concept (Figure 6).⁶⁸ The Dunitz-Waser concept was developed for cycloalkenes and has postulated that a *trans* double bond can be substitute by a long bond and a *cis* double bond by a phantom atom to understand the dynamics of cycloalkenes. This concept was translated into cyclic peptides by considering a peptide bond equivalent to a C=C double bond.⁶⁹

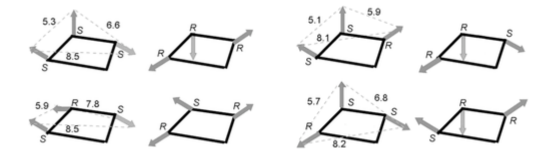


Figure 6 Schematic Dunitz–Waser topographic depiction of the eight PR-CTP models (enantiomeric pairs), showing the *S*/*R* stereochemistry of the C α , and the distances (° A) between the C β atoms. Gray arrows indicate the pseudo-axial or pseudo-equatorial disposition of the side chains (C α -C β vectors).

All stereoisomers show a strong preference for a secondary structure characterized by a β -turn centered on β -Phe- ψ (NHCO)Ala, and a second one on Phe- β -Phe- ψ (NHCO).⁷⁰ This feature correlates to the strong tendency of the diamine amide protons to participate to H-bonded structures. This observation is not obvious;

indeed, literature precedent suggested that configurational changes in a cyclopeptide can lead to alternative secondary structures.^{37,40,60,62,63} Besides, it can be appreciated that the stereoisomeric scaffolds actually differ in the directions of the C α -C β vectors of Ala, diamine, and L/D-Phe.

The eight stereoisomers of the PR-CTP sequence c[β -Phe- ψ (NHCO)Asp- ψ (NHCO)Gly-Arg] containing 3phenylpropane-1,2-diamine (a β -Phe retro-mimic), and a malonate (a retro-Gly), were synthesized. The ability of the resulting compounds to inhibit the adhesion of the α 5 β 1 integrin-expressing K562 cells (human erythroleukemic cells) or $\alpha\nu\beta$ 3 integrin-expressing SK-MEL-24 cells to immobilized FN, as well as the adhesion of $\alpha\nu\beta$ 3 integrin-expressing HUVEC cells (human umbilical vein endothelial cells) to immobilized VN, was evaluated. Compound **15**, c[(*R*)- β -Phe- ψ (NHCO)Asp- ψ (NHCO)Gly-Arg] (Figure 7), containing (*S*)-Arg and (*S*)-Asp, exhibited the highest potency as an inhibitor of cell adhesion mediated by $\alpha\nu\beta$ 3 and α 5 β 1 integrins.⁷¹ The observed IC₅₀ value for α 5 β 1 integrin/FN (K562 cells) was 2.4×10⁻⁸ M, that is about 10 times lower than that exhibited against $\alpha\nu\beta$ 3 integrin, IC₅₀ $\alpha\nu\beta$ 3/FN (SK-MEL-24 cells) = 1.8×10⁻⁷ M, IC₅₀ $\alpha\nu\beta$ 3/VN (HUVEC cells) = 2.4×10⁻⁷ M. The diastereoisomer **16**, c[(*S*)- β -Phe- ψ -(NHCO)Asp- ψ (NHCO)Gly-Arg] (Figure 7), differing from **15** exclusively by the configuration of the diamine moiety, maintained sufficient affinity for α 5 β 1 integrin, but increased selectivity against $\alpha\nu\beta$ 3 integrin Indeed, although **16** exhibited an IC₅₀ $\alpha\nu\beta$ 3/VN = 2.5×10⁻⁵ M.

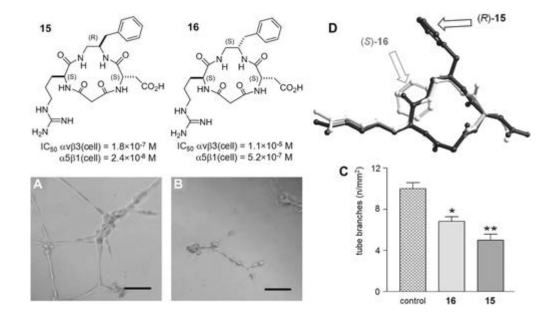


Figure 7 Structures of the PR-CTP RGD mimetics **15** and **16**; IC50 values refer to integrin-expressing cells (cell). (**A** and **B**) The $\alpha 5\beta 1/\alpha v\beta 3$ mixed integrin antagonist **15** inhibits bFGF-induced formation of capillary-like tubes of HUVEC on Matrigel; (**A**) HUVEC plated on Matrigel in the presence of bFGF; (**B**) HUVEC treated with compound **15** (1 μ M), black bar = 100 μ m. (**C**) Mean numbers of tubes per square millimeter ± SEM; **P* < 0.01, ***P* < 0.05 vs. control (Newman–Keuls test after ANOVA). (**D**) Overlap of the backbone structures of **15** (black) and **16** (gray), determined by ROESY and molecular dynamics.⁷¹

The potential antiangiogenic activity of these novel antagonists of the integrins $\alpha 5\beta 1$ and $\alpha \nu \beta 3$, was estimated *in vitro* by analyzing the formation of capillary tubes composed of HUVEC cells cultured in Matrigel, when stimulated by basic Fibroblast Growth Factor (bFGF). HUVEC cells plated in the presence of bFGF alone gave rise to a nice capillary-like network (Figure 7A). The bFGF-stimulated formation of the capillaries was significantly inhibited by compounds **15** and **16** (for compound **15**, see Figure 7B). The number of tube branches per square millimeter was reduced from 10 ± 3 (bFGF-treated HUVEC) to 5.1 ± 0.7 by treatment with 1 μ M **15** and 6.8 ± 0.9 with 1 μ M **15** (Figure 7C).

The biological activities of **15** and **16** appear somewhat contrasting, despite the fact that the amino acid sequence of the peptides is identical. Conformational analysis was performed and confirmed the side chains 3D display as predicted from the corresponding 13-membered PR-CTPs models (Figure 6). The two stereoisomers share the same disposition of the RGD-mimetic region. The average distance between the β carbons of Asp and Arg, (~8.5 Å), and the distance between the carboxylate and guanidino groups (~13.5 Å), are very similar to that found in the α 5 β 1 or α v β 3 selective RGD-cyclopeptides described in Section 2. The most relevant 3D difference can be noticed in the diamine region; apart from the different position of diamine amide protons, the benzyl side chain of the diamine is nearly equatorial in **15**, while it adopts an opposite axial position in **16** (Figure 7D).

The different $\alpha\nu\beta3$ receptor affinities were rationalized with the aid of docking studies performed on the model of binding site derived from the crystallographic structure of the extracellular segment of integrin $\alpha\nu\beta3$ in complex with cilengitide **3** (PDB ID: 1L5G).⁴⁵ The combination of conformational analyses and molecular docking support the fundamental role of the aromatic ring adjacent to Asp in determining the different pharmacological profile. Indeed, the pseudo-equatorial benzyl group in **15** points toward the outside of the $\alpha\nu\beta3$ integrin binding site, preventing any steric hindrance, and reproducing the orientation present in cilengitide. By contrast, in compound **16** the axial position of the aromatic side

chain is responsible for an unfavorable steric interaction that prevents the compound from entering into the receptor site (Figure 8).

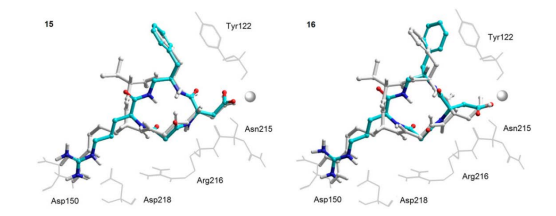


Figure 8 Sketches of the structures of **15** and **16**71 determined by molecular docking (Glyde version 4.5) of the ligands within the crystal structure of the integrin $\alpha v \beta 3$ (PDB ID: 1L5G).⁴⁵

As for the integrin $\alpha 5\beta 1$, at that time the 3D crystal structure was not available yet, and few details about ligand-receptor interactions were reported in the literature. However, a homology model of the receptor⁴⁷ was developed and utilized to design selective $\alpha 5\beta 1$ integrin ligands.⁴⁸ On the basis of this model, it was postulated that the integrin $\alpha 5\beta 1$ could host the benzyl side chain of both compounds. Indeed, the comparatively larger binding site of $\alpha 5\beta 1$ integrin compared to that of integrin $\alpha v\beta 3$ can easily accommodate the axial phenyl ring of **16**, leading to a pronounced selectivity for $\alpha 5\beta 1$ over $\alpha v\beta 3$ integrin, while the equatorial disposition of **15** can fit both integrins.

The better $\alpha v\beta 3$ integrin receptor affinity of (*S*,*R*,*S*)-PR-CTP **15** respect to the (*S*,*S*,*S*)-configured **16** nicely confirms the predictive efficacy of the PR-CTP topographic models discussed above. The positioning and the distances between C β were correlated to the distances found in the bioactive conformation of the prototypic $\alpha v\beta 3$ -selective cilengitide **3**, discussed in the previous sections (Figure 9A). The comparison highlighted the good match between the topographic Dunitz-Waser depictions of **3** (Figure 9B) and of the (*S*,*R*,*S*)-CTP model (Figure 9C), and the poor overlap with the sketch of (*S*,*S*,*S*)-CTP concerning the benzyl side chain (Figure 9C).

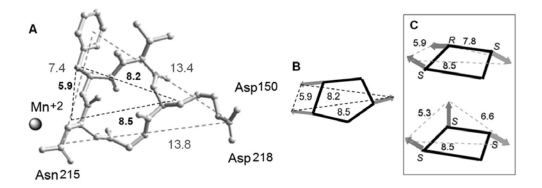


Figure 9 (A) Bioactive conformation of the $\alpha\nu\beta$ 3-selective c[RGDf(*N*-Me)V], cilengitide 3, showing the distances (A°) between the pharmacophores (gray digits and gray dashed lines) and between C β atoms of Asp, D-Phe, and Arg (black digits and black dashed lines). (B) Dunitz–Waser sketch of 3. (C) Dunitz–Waser sketches of (*S*,*R*,*S*)-CTP (top), and (*S*,*S*,*S*)-CTP (bottom), models of 15 and 16, respectively.

3.2. Ligands with external cyclic scaffolds of β-amino acid structure/span

A few years ago, Sewald and coworkers reported the synthesis, the pharmacological evaluation, and structural determination, of several novel ligands for integrin $\alpha\nu\beta3$, including the pentapeptides c[Arg-Gly-Asp-(+)- β -Acc-Val] (17) and c[Arg-Gly-Asp-(-)- β -Acc-Val] (18) containing enantiomeric (+)- or (-)-*cis*- β -amino cyclopropanecarboxylic acid (β -Acc) adjacent to the RGD motif.⁷² The biological activity of peptides 17 and 18 was determined in adhesion experiments with cancer cell lines. Peptide 17, containing (+)- β -Acc, displayed a ten-fold higher affinity (IC₅₀ = 2.0×10⁻⁸ M) than the reference peptide c[RGDfV] 2, as determined by adhesion experiments of WM115 cells to VN. By contrast, the inhibitory effect of the diastereomeric 18 was comparable to that of 2. The adhesion of K562 cells mediated by α 5 β 1 integrin to FN was inhibited to nearly the same extent by peptides 17 and 18, with IC₅₀ values of 1.5×10⁻⁶ M and 1.8×10⁻⁶ M, respectively; that is, more active than reference peptide 2.

The significantly different affinity of the β -Acc-RGD peptides to $\alpha\nu\beta3$ integrin can be interpreted as a consequence of the stereochemistry of the β -Acc residue. Conformational analysis revealed that in **17** glycine is located in the central position of a γ -turn, while the (+)- β -Acc residue occupies the *i*+*1* position of a *pseudo* β -turn (Figure 10). In peptide **18**, Asp is in the central position of an inverse γ -turn. Between the amino acids (-)- β -Acc and Gly, the structure adopts the torsional angles characteristic of a 3¹⁰ helix, thus corresponding to a conformation formerly known as the β III turn with value in the *i*+*1* position and arginine in the *i*+*2* position

(Figure 10). All this accounted for a more extended geometry of the RGD sequence for **18** relative to **17**, so the latter is more compatible with the structural requirements for optimal binding to integrin $\alpha\nu\beta3$ (Section 2), namely a short distance between the C β atoms of Arg and Asp of about 6.5-8.5 Å.

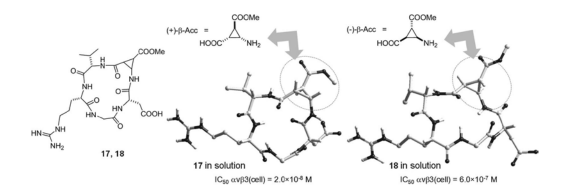


Figure 10 Structures of the (+)- and (-)- β -Acc, and β -Acc-containing peptides **17** and **18**. Lowest-energy conformations obtained by restrained MD calculations based on experimental C-C bond distance information. IC50 values refer to integrin-expressing cells (cell).

Albericio, et al. reported the efficient solid phase preparation of cyclic RGD peptides, connected 'head-totail' by the side chains of a diketopiperazine (DKP) rigid scaffold, giving rise to a new class of bicyclic RGD peptides. The DKP scaffold was obtained in solid phase by cyclization of a dipeptide orthogonally protected at the side chains, H-Lys(N^{ϵ} -Mtt)-Asp(β Allyl)-OMe, attached to a Backbone Amide Linker (BAL) resin by the peptide bond nitrogen. The affinity of the RGD-containing ligands comprising five to seven residues towards the $\alpha\nu\beta3$ integrin was determined in binding assays using the radioligand [¹²⁵I]echistatin. Results indicated that the compound which contained just the sequence RGD and the scaffold showed the highest binding affinity.⁷³

In this context, some years later Piarulli, et al. developed a similar bifunctional DKP scaffold. The sixmembered heterocyclic ring was prepared from (*S*)-2,3-diaminopropionic acid and L-aspartic acid (Figure 11); therefore, this scaffold includes two vicinal β -amino acids, namely a α , β^2 -diaminoacid and a β -carboxy- β^3 amino acid.

The RGD sequence was coupled to either *cis*-DKP or *trans*-DKP (Figure 11). Following macrolactamization, the resulting cyclic RGD peptidomimetics **19** (*cis*-DKP) and **20** (*trans*-DKP) were examined *in vitro* as inhibitors of biotinylated VN binding to $\alpha\nu\beta3$ and $\alpha\nu\beta5$ receptors.⁷⁴ Interestingly, **20** exhibited a nanomolar affinity for the $\alpha\nu\beta3$ integrin, IC₅₀ = 3.2×10^{-9} M, and 40-fold selectivity over $\alpha\nu\beta5$,

while **19** showed only a modest micromolar affinity for the $\alpha\nu\beta3$ integrin and no activity towards $\alpha\nu\beta5$ integrins, IC₅₀ = 3.9×10^{-6} M.

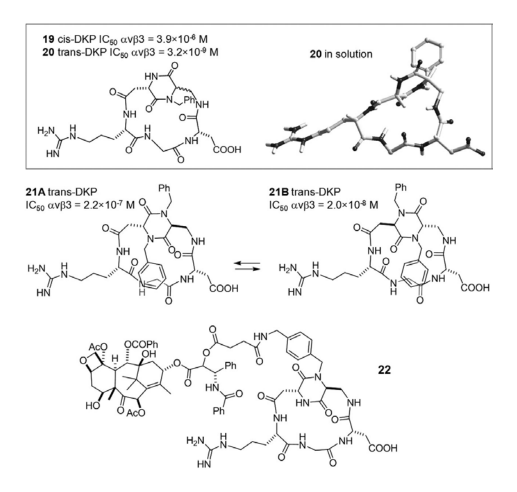


Figure 11 In the box: structures of cyclic RGD-peptidomimetics **19** and **20** containing diketopiperazine scaffolds, and 3D structure of **20** as obtained by restrained Monte Carlo/Stochastic Dynamics (MC/SD) simulations based on experimental bond distance information, showing a pseudo β -turn at DKP-Arg. Diastereomers **21A** and **21B**, resulting from hindered rotation. Structure of the c[DKP-RGD]-paclitaxel conjugate **22**. IC50 values refer to isolated integrins.

The different performance of the DKP-RGD-mimetics **19** and **20** was investigated by NMR spectroscopy and computational methods. Conformational analysis revealed that **19** exists as two interconverting conformers; NOESY spectrum showed two mutually exclusive long-range NOE contacts, one indicative of a conformer characterized by a β -turn centered on Gly-Asp, and a second one showing a β -turn centered on Arg-Gly. Conversely, the conformation of the RGD-mimetic **20** containing the *trans*-DKP is rather rigid, with a clear β -turn on DKP-Arg stabilized by a H-bond between Gly-NH and C5=O (Figure 11), and a distance between the C β of Arg and Asp of 9.4 Å. Docking computations of this extended conformation of **20** afforded a lowest-energy conformer that preserves the ionic interactions of cilengitide **3** in complex with integrin $\alpha v \beta 3.^{45}$ By contrast, the docked geometries of **19** produced top-ranked poses conserving optimal interactions only with the α subunit.

Subsequently, the Gennari group expanded the library of RGD mimetics by introducing the DKP scaffolds of differing configuration at the two stereocenters, as well as different *N*-substitution patterns. Eight resulting cyclopeptides were analyzed as inhibitors of the binding of biotinylated VN to integrins $\alpha\nu\beta3$ and $\alpha\nu\beta5$. The peptidomimetics derived from *trans*-DKPs gave nanomolar IC₅₀ values.⁷⁵ Interestingly, the 2:1 **21A:21B** diastereomeric/rotamer pair was separated owing to hindered rotation of the dibenzyl-DKP scaffold (Figure 11). Poor receptor affinity was obtained for ligand **21A**, IC₅₀ $\alpha\nu\beta3 = 2.2 \times 10^{-7}$ M, while rotamer **21B** was the most potent ligand of the series, IC₅₀ $\alpha\nu\beta3 = 2.0 \times 10^{-8}$ M).

It was established that rotamer **21B** adopts a variety of conformations in solution, including an H-bonded type I β -turn having Gly-Asp at the positions *i*+1, *i*+2, characterized by an extended RGD sequence. Docking calculations starting from this conformation gave top-ranked binding modes consistent to the bioactive conformation of cilengitide **3** within the integrin (see above).⁴⁵

In order to investigate the binding epitope of the c[DKP-RGD] peptidomimetics to the integrins, NMR studies based on saturation transfer and detection of transferred NOEs were used for analyzing the ligands during their interaction with living cells known to express on their membrane the integrins $\alpha\nu\beta3$ (ECV304 bladder cancer cells) and α IIb $\beta3$ (human platelets).⁷⁶ This technique was originally introduced by Meyer for applications in the integrin field.⁷⁷ Transferred NOEs can be recorded only when there is rapid exchange, this meaning that the binding should not be too high and only ligands with relative low binding affinities can be investigated. The Meyer group therefore used the integrin α IIb $\beta3$ and a $\alpha\nu\beta3$ selective ligand with much lower affinity to the investigated integrin. The NMR results were supported by docking studies and were compared to the results of competitive $\alpha\nu\beta3$ receptor binding assays and competitive ECV304 cell adhesion experiments, and clearly confirmed that only those ligands with *trans*-RGD fragments are able to interact with both the α and β integrin subunits via an electrostatic clamp.

The c[DKP-RGD] ligands were exploited for integrin-mediated delivery of the cytotoxic drug paclitaxel (PTX).⁷⁸ The c[DKP-RGD]-PTX constructs were good integrin ligands *in vitro*, since they inhibited

biotinylated VN binding to the purified $\alpha\nu\beta3$ integrin receptor at low nanomolar concentration. Furthermore, when tested against a panel of human tumor cell lines (including the cisplatin-resistant IGROV-1/Pt1), they showed cytotoxic activity with efficiency similar to that of paclitaxel. In particular, the c[DKP-RGD]-PTX **22** (Figure 11) emerged as a candidate for *in vivo* testing, due to its good activity and stability under physiological conditions and in plasma. Compound **22** exhibited higher tumor-targeting activity compared to PTX alone against an IGROV-1/Pt1 human ovarian carcinoma xenotransplanted in nude mice.

4. Integrin ligands with β-amino acid residues in place of Asp

One of the larger β 3 integrin antagonist classes employs β -amino acids to mimic the aspartate residue of the RGD mimetic. Structure-activity relationship (SAR) investigations have revealed the potent activity of agents which have substituents appended to both the α and β position of the β -amino acid units of these antagonists. Several clinical candidates targeting platelet α IIb β 3 and containing these β -amino acid units were evaluated clinically. Since the many integrin peptidomimetic ligands of this kind have been previously reviewed exhaustively,^{23,38,79} the following sections report selected representative examples from the recent literature.

4.1. Recent examples of β-amino acid analogues of Asp, and their uses in medicinal chemistry

Simple β -alanines⁸⁰ or β -alanine derivatives carrying different substituents at the β position have been successfully applied in the synthesis of RGD mimetics.^{79,81,82} Most ligands include aromatic substituents to mimic an Asp-Phe dipeptide,^{83,84} but also other groups have been utilized (e.g. an alkyne⁸⁵).

By far, the most widely used β -amino acid equivalent is diaminopropionate.^{23,38,79} Very often, the α -amino group is utilized to introduce diverse types of substituents, e.g. aryl, sulfonyl,⁸⁶ arylsulfonyl⁸⁷ groups etc., in close proximity to the carboxylic acid pharmacophore. These aromatic groups were shown to contribute to receptor affinity thanks to an additional, energetically favorable "exosite" π - π interaction with Tyr178 of αv .⁸⁸

One extraordinary advantage of diaminopropionate is the opportunity that it provides to conjugate the integrin ligand to nanoparticles, antibodies, lipids, fluorescent tags, among others,^{27,28} or for oligomerization

of the ligand.⁸⁹ For instance, the diaminopropionate segment allowed conjugating an oligomeric peptidomimetic integrin $\alpha\nu\beta$ 3 antagonist to a monoclonal antibody (MoAb), in order to increase integrin $\alpha\nu\beta$ 3 receptor-binding avidity (Figure 12). The antibody moiety served as carrier of the antagonist **23** to target tumor-induced neovasculature.⁹⁰

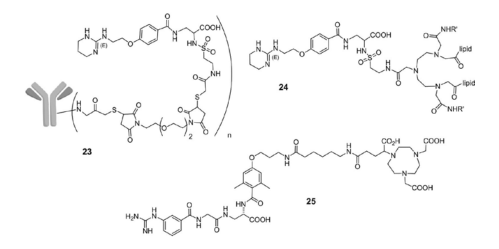


Figure 12 Structures of an integrin antagonist-conjugated MoAb 23, a lipid-conjugated integrin antagonist 24, and a NODAGA-conjugated integrin antagonist 25.

The same $\alpha v\beta$ 3-targeting integrin ligand was coupled through the diaminopropionate unit to branched lipids **24** capable to form a cationic nanoparticle (Figure 12), in order to selectively deliver genes to angiogenic blood vessels in tumor-bearing mice.⁹¹

Kessler et al. functionalized a α 5 β 1-selective antagonist (Figure 12) with 1-((1,3-dicarboxy)propyl)-4,7-(carboxymethyl)-1,4,7-triazacyclononane (NODAGA).⁹² This strategy provided the compound **25**, an effective and selective α 5 β 1-integrin antagonist that enabled specific molecular imaging *in vivo* by PET, based on the different integrin pattern of tumor cells.

The side-chain amino group of diaminopropionate was also functionalized to append biotin or a biotin derivative containing an oligo(ethylene glycol) ((EG)4) linker, to give a bifunctional RGD-based peptidomimetic particularly effective for surface presentation; the surfaces decorated with this ligand promoted both cellular adhesion and integrin activation.⁹³

4.2. Retro-sequences containing isoAsp residues

Some researchers proposed that the spontaneous modification of the Asn-Gly-Arg (NGR) motif in FN ligands into *iso*DGR might result in a gain of integrins adhesion by creating a new binding site.^{94,95} This hypothesis was critically examined using recombinant FN with mutations in the NGR motif.⁹⁶ Subsequent studies confirmed that the *iso*DGR sequence maintains the same binding mode of RGD, affording the classic electrostatic clamp, as well as the normal additional polar interactions.^{97,98}

In this field, Kessler, et al. grafted the *iso*DGR motif onto the structures of retro-RGD cyclopentapeptides. The use of an *iso*Asp residue does not modify the span between the guanidinic and carboxylate pharmacophores; however, it provokes an enlargement of the macrocyclic peptide similar to that caused by the introduction of one β -amino acid (see Section 3.1). Specifically, from 15 bonds present in the normal cyclopentapeptides, the substitution leads to 16 bonds present in the *iso*DGR analogues. Therefore, this structural modification is expected to result in considerable conformational bias of cyclopentapeptide backbones.

On the other hand, binding experiments showed that the sequential position of the aromatic amino acid and glycine flanking *iso*DGR, determines the affinities of the pentapeptides to either integrin $\alpha\nu\beta3$ or $\alpha5\beta1$. Indeed, c[phg*iso*DGRG] **26** (phg = D-phenylglycine) showed enhancement of pharmacological activity towards $\alpha5\beta1$ integrin, IC₅₀ $\alpha5\beta1 = 1.9 \times 10^{-8}$ M as determined by competitive ELISA using immobilized ECM protein and soluble integrin, relative to integrin $\alpha\nu\beta3$ (IC₅₀ > 10⁻⁶ M).⁹⁹ By contrast, c[G*iso*DGRphg] **27** was more active towards integrins $\alpha\nu\beta3$ (IC₅₀ = 8.9×10^{-8} M) (Figure 13). Molecular docking were performed for the $\alpha\nu\beta3$ receptor, using the crystal structure of the cilengitide- $\alpha\nu\beta3$ integrin complex, and the $\alpha5\beta1$ receptor, starting from the homology model seen in Section 2.⁴⁵ The simulations confirmed the usual interaction patterns. Furthermore, selectivity toward FN-binding integrins $\alpha\beta31$ and $\alpha\nu\beta6$ relative to VN-binding integrins $\alpha\nu\beta3$ and $\alpha\nu\beta5$ was improved by appropriate substitution of the flanking amino acids in the peptide.¹⁰⁰

Very recently, Gennari, et al. synthesized the cyclopeptides **28** and **29**, comprising *iso*DGR (Figure 13) and the bifunctional DKP scaffolds,^{74,75} and determined their ability to compete with biotinylated VN for binding to the purified $\alpha\nu\beta3$ and $\alpha\nu\beta5$ receptors. In particular, compound **29** showed the lowest IC₅₀ recorded value for integrins $\alpha\nu\beta3$ (IC₅₀ = 9.2×10⁻⁹ M), and good selectivity over integrin $\alpha5\beta1$.¹⁰¹

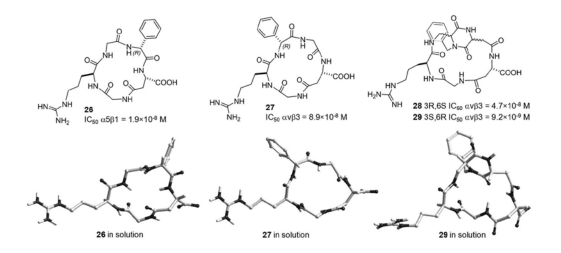


Figure 13 Structures of cyclic pentapeptides **26**, **27** containing the *iso*DGR motif, and of the cyclic *iso*DGR peptidomimetics **28** and **29** containing the bifunctional DKP scaffolds; 3D molecular structure of **26**, **27**,99,100 and *trans-(3S,6R)-DKP-iso*DGR **29**101 obtained by restrained MC/SD simulations based on experimental bond distance information, followed by energy minimization. IC50 values refer to isolated integrins

Data obtained from NMR experiments demonstrated the high conformational flexibility of compound **28**, while in the case of peptide **29**, experiments suggested the existence of two preferred conformations, a β -turn with a H-bond between DKP-NH10 and *iso*AspC=O, and a *pseudo* β -turn conformer stabilized by an H-bond involving Gly-NH and DKP-C5=O (Figure 13). Docking calculations confirmed that the latter structure corresponds to the bioactive conformation.

Finally, the *iso*Asp residue was also utilized both in straight or retro sequences by Gentilucci et al. in the preparation of conformationally constrained mimetics of the $\alpha 4\beta 1$ integrin antagonist BIO1211 (see Section 5.2.2).¹⁰²

5. Integrin ligands containing β -amino acid cores

A variety of cyclic (aromatic and non-aromatic) or acyclic scaffolds, with chiral and achiral structures, was examined to furnish Gly surrogates in RGD peptides.^{22,23,38} Generally, only flat structures within the RGD mimetic permit to maintain the effective interactions with integrin receptors. Sugar and piperazine scaffolds which have shown poor or modest anti- $\alpha v\beta 3$ activity, usually present almost flat structures, due to the all-equatorial conformations.^{103,104,105,106} By contrast, replacement of the Gly residue by Ala, bulky, longer, or branched core residues, resulted in a dramatic loss in potency.^{38,107} The introduction of β -amino acids or other β -amino acid derived moieties is expected to provide hybrid molecules with increased stability against

enzymatic hydrolysis. Clearly, the increased length of the β -amino acid relative to Gly impacts not only the distance between the pharmacophore-containing moieties, but also the conformations of the RGD tripeptide motif. For this reason, only peptidomimetic compounds specifically designed to maintain the correct shape and distances can successfully include a β -amino acid within the RGD sequence.

5.1. Integrin ligands containing acyclic β-amino acid cores

Some years ago, Hayashi et al. reported the potent non-peptidic α IIb β 3 integrin antagonist **30** (NSL-95301), with a unique trisubstituted β -amino acid unit utilized as central core. The β -amino acids were prepared from the corresponding β -lactams (Figure 14), prepared in turn by a convenient ester-imine condensation. Following conformational analyses and SAR studies, the optimization of the amidine group led to the potent and orally active antagonist of the platelet FB receptor **31** (Figure 14), containing 3-ethyl-2,2-dimethyl- β -alanine. The active (*R*)-enantiomer of **31** effectively inhibits platelet aggregation *in vitro*.¹⁰⁸ Tested *in vivo* in guinea pigs, **31** was orally active; the onset of antiplatelet action was rapid, albeit the duration was relatively brief.

The trisubstituted β -residue controls the active conformation of the molecule. Molecular modeling of **30** and related compounds revealed a quite rigid albeit linear structure, which effectively fixes the distance between the pharmacophoric ends. The amidino and the carboxyl groups project at opposite sides of a cup-shaped structure, the phenyl group at the β -position being perpendicular to the piperidine ring and amidinophenyl group. This cup-shaped rigid conformation originates from the correlated and restricted motions of the piperidine ring, the *geminal* groups at the α -position, and the β -substituent; the latter appeared to be very important to restrict the overall flexibility.

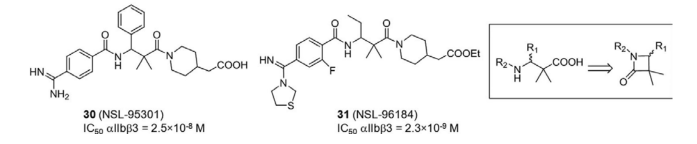


Figure 14 Structures of the α IIb β 3 integrin antagonists **30** and **31** containing the trisubstituted β -residues; in the box: retrosynthesis of the β -residues.

More recently, Cardillo, et al. reported the enantioselective transesterification or acylation of racemic β amino esters by treatment with *Candida antarctica Lipase B* (CaLB)-catalyzed, for the isolation of enantiopure intermediates of pharmacological interest **32** and **33** (Figure 15). Both the malonyl and the β -amino esters proved to be suitable substrates for CaLB. The enzyme recognized under kinetic resolution conditions the less hindered position, and the transesterification reaction proceeded with good yields and high enantioselectivity. This strategy was applied to the two-step synthesis of the RGD mimetic compound **34**,¹⁰⁹ which showed moderate ability as inhibitor of cell adhesion mediated by integrins $\alpha\nu\beta$ 3 and α 5 β 1, with IC₅₀ values of 1.7×10⁻⁷ M and 9.7×10⁻⁷ M, respectively.¹¹⁰

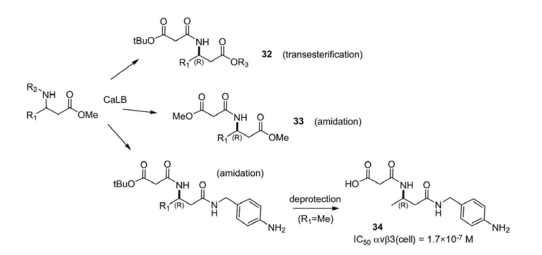


Figure 15 General strategies for obtaining sequences **32**, **33** containing enantiopure β -amino acids, including the RGD mimetic **34**, by CaLB-catalyzed resolution of racemic β -amino acid derivatives. IC50 value refers to integrin-expressing cells (cell).

Aiming at obtaining small mimetics of the RGD motif, Cardillo et al. introduced dehydro-β-amino acid as central scaffolds instead of Gly. Enantiopure allylic alcohols were prepared by an enzymatic approach, and the allylic amination of the deriving carbonates¹¹¹ with 4-substituted benzylamines proceeded through an SN2' mechanism (Figure 16A), giving access to a novel type of RGD mimetic requiring few protection-deprotection extra steps.¹¹²

Starting from racemic carbonates, Tolomelli, et al. prepared various RGD mimetic candidates and tested them in the inhibition of $\alpha\nu\beta3$ and $\alpha5\beta1$ integrin-mediated cell adhesion.¹¹³ The short molecules **35a-d** (Figure 16B) showed unexpected activity, and selectivity correlated to the substituent at position 3. Derivative **35a** (R = Me) did not affect $\alpha\nu\beta3$ integrin-mediated cell adhesion, while it showed excellent inhibition of the $\alpha5\beta1$ integrin (IC₅₀ = 1.9×10⁻⁸ M). On the other hand, the compounds **35b** (R = *i*Pr) and **35c** (R = thiophenyl) had micromolar affinity toward both integrins, while **35d** (R = cyclohexyl) showed preferential affinity towards $\alpha\nu\beta3$ receptor. Nevertheless, the study of additional peptidomimetics is required before a clear understanding of SAR is reached. Moreover, these molecules are quite small relative to the usual RGD sequences, therefore the different side chains may be less determinant for integrin affinity. Few antagonists of similar length have been reported in the literature; for instance, the heterocyclic α IIb $\beta3$ antagonist RUC-1 was shown crystallographically to bind only to the α IIb subunit, in contrast to classic RGD antagonists.¹¹⁴ Docking analysis aimed at determining the binding mode of the short compound **35b** with the integrin $\alpha\nu\beta3$ were inconclusive. Indeed, the computations gave alternative plausible poses characterized by very similar docking scores, one showing ionic interactions with both the α (MIDAS) and β (Asp²¹⁸) subunits, plus a stacking interaction between the phenyl group of the ligand and Tyr¹⁷⁸, and a second pose in which the ligands maintain only the electrostatic interaction with MIDAS in the α vubunit.¹¹⁵

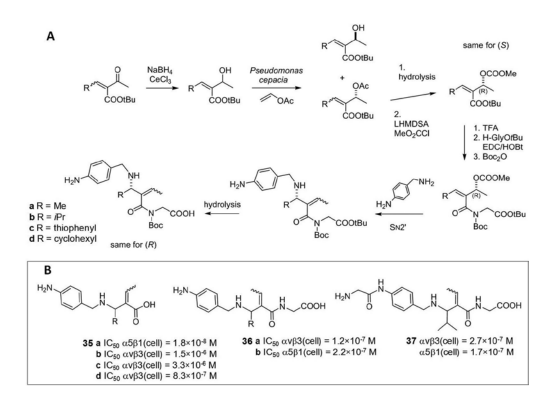


Figure 16 (A) Synthesis of enantiomerically pure allylic carbonates and synthesis of dehydro- β -amino esters through amination of allylic carbonates. (B) In the box: ligands 35–37 containing dehydro- β -amino acid scaffolds, examined for integrins $\alpha\nu\beta3$ and/or $\alpha5\beta1$. IC50 values refer to integrin-expressing cells (cell).

The larger ligand **36b** with R = iPr (Figure 16) exhibited an increased $\alpha 5\beta 1$ integrin-mediated inhibitory activity (IC₅₀ $\alpha 5\beta 1 = 2.2 \times 10^{-7}$ M; IC₅₀ $\alpha v\beta 3 = 2.0 \times 10^{-6}$ M) in comparison to the shorter analog **35b**. By contrast, compound **36a** with R = Me displayed good activity and a strong preference toward $\alpha v\beta 3$ integrin (IC₅₀ = 1.2×10^{-7} M). The molecule was further elongated by introduction of a second glycine unit (Figure 16); this modification afforded the most potent dual antagonist **37** (IC₅₀ $\alpha v\beta 3 = 2.7 \times 10^{-7}$ M; IC₅₀ $\alpha 5\beta 1 = 1.7 \times 10^{-7}$ M).

The effect of the most effective compounds **36a** and **37** on intracellular $\alpha\nu\beta$ 3 integrin-mediated signaling activation was assessed using DAOY medulloblastoma cells. Pre-incubation with both compounds produced a measurable decrease of the phosphorylation of extracellular-signal-regulated kinases (ERK) induced by FN. In addition, the effect of both compounds on intracellular α 5 β 1 integrin-mediated signaling activation was analyzed using K562 cells. Compound **37** reverted FN-induced phosphorylation of ERK1/2. By contrast, compound **36a** was not able to reduce ERK pathway activation, which is suggestive of a different mechanism of action.

The different performances of **36b** and **37** in $\alpha\nu\beta$ 3-mediated cell adhesion inhibition were analyzed by docking computations, using the X-ray crystal structure of the ectodomain of integrin $\alpha\nu\beta$ 3 complexed with the cyclic pentapeptide ligand cilengitide **3** (Section 2).⁴⁵ The pose of compound (*3R*,*Z*)-**36b** with the best docking score shows the carboxylate group coordinated to the divalent cation in the MIDAS region of the β 3 subunit, while the ligand's aromatic amine interacts with the negatively charged side chain of Asp²¹⁸ in the $\alpha\nu$ subunit. Nevertheless, for the latter moiety the fit with the cilengitide Arg guanidinium group is not precise. As for compound (*3R*,*Z*)-**37**, the introduction of an additional glycine gave the correct distance required for a good fit with the receptor. The best calculated pose of (*3R*,*Z*)-**37** showed the acid and basic pharmacophoric groups involved in strong ionic bonds with charged regions of the receptor binding site, nicely reproducing the electrostatic clamp of the RGD sequence of integrin-bound **3**.

5.2. Integrin ligands containing heterocyclic β-amino acid cores

One approach to obtain highly stable and structurally defined peptidomimetics is to incorporate a conformationally restricted residue, introducing a local constraint which may result in a significant reduction of the accessible conformational space.¹¹⁶ In this context, a small-to-medium sized heterocycle can be incorporated as central core of a peptide, to orient the pharmacophoric side chains and to fix the adequate distanced for binding. Generally, heterocyclic scaffolds of minimal structural complexity and steric requirements have been designed to prevent undesired interactions. The following sections are dedicated to illustrative examples of integrin ligands containing a heterocyclic core having a β -amino acid structure. Representative experimental protocols are included to highlight salient synthetic procedures for the preparation of non-racemic scaffolds.

5.2.1. Integrin ligands containing β-lactam scaffolds

Claudio Palomo et al. proposed the use of peptides containing β -lactam variants of the classic Freidinger lactams¹¹⁷ as conformationally restricted nucleators of β -turn structures. Starting from these scaffolds, specific cyclic RGD β -lactam peptidomimetics were also developed.¹¹⁸

Very recently, Galletti et al.¹¹⁹ designed novel integrin ligands including a β -lactam scaffold, which can be regarded as an *N*-to-*C* β cyclic β^3 -amino acid (Figure 17). The compounds **38** and **39** were readily prepared from the commercially available 4-acetoxy-azetidin-2-one. The carboxylic acid function at the C4 side chain of the β -lactam was introduction by Reformatsky reaction, followed by *N*-acylation to give the RGD mimetic **38**. The insertion of a further Gly at the C4 side chain afforded the longer compound **39**.

These new derivatives were tested in their ability to modulate the adhesion of integrin-expressing cells to immobilized FN. Surprisingly, the collected data support that the β -lactam derivatives present an interesting activity as $\alpha\nu\beta3$ and $\alpha5\beta1$ integrin agonists. Indeed, **38** and **39** increased in a concentration-dependent manner the FN-mediated adhesion of K562 cells and SK-MEL-24 cells. In particular, ligand **38** exhibited higher affinity towards $\alpha5\beta1$ integrin (EC₅₀ = 1.2×10^{-8} M), while **39** was selective for integrin $\alpha\nu\beta3$ (EC₅₀ = 1.1×10^{-8} M). Both ligands were unable (IC₅₀ > 1×10^{-4} M) to modify the adhesion of Jurkat E6.1 cells to vascular cell

adhesion molecule-1 (VCAM-1) mediated by $\alpha 4\beta 1$ integrin, and the adhesion to intercellular adhesion molecule-1 (ICAM-1) mediated by $\alpha L\beta 2$ integrin.

The new β -lactam derivatives maintained their ability to increase cell adhesion even in the absence of FN. In a second series of experiments, the wells of a standard test plate were coated by passive adsorption with either **38**, **39**, FN as a positive control, bovine serum albumin (BSA) as a negative control. K562 and SK-MEL-24 cells were seeded for adhesion; the two cell lines exhibited significant adhesion to both ligands, comparable to FN, while the cells did not adhere to wells coated with BSA. Antibodies of the β 1 or α v integrin subunit added in advance to the cells blocked the adhesion mediated by **38** and **39**, strengthening the hypothesis that cell adhesion was mediated by the integrins $\alpha\nu\beta$ 3 and $\alpha5\beta$ 1. Interestingly, the adhesion of K562 cells to both ligands was partially blocked by the selective α IIb β 3 integrin antagonist tirofiban. On the basis of these data, the authors conjectured that both $\alpha5\beta$ 1 and α IIb β 3 integrins may contribute to mediate the adhesion of K562 cells induced by **38** and **39**.

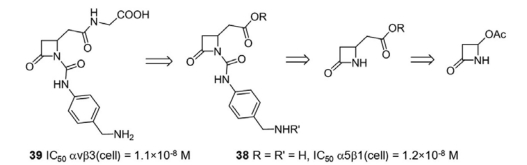


Figure 17 Retrosynthetic strategy for the β -lactam-containing RGD-like compounds **38** and **39**. IC50 values refer to integrin-expressing cells (cell).

5.2.2. Five-membered heterocyclic β-amino acidic cores

Dominguez et al. proposed a conformationally constrained *N*-aryl- γ -lactam scaffold to mimic the Arg-Gly dipeptide. This γ -lactam structurally resembled an *N*-to-C α cyclic β^2 -residue, and was obtained from nitroaniline and itaconic acid (Figure 18). The pure enantiomers of the γ -lactams were obtained via chemical resolution by means of the Evans' chiral auxiliary. When elaborated with various β -amino acids, the resulting ligands provided potent and selective $\alpha\nu\beta3$ antagonists. All compounds were assayed in competitive

electrochemiluminescent binding experiments using VN as specific $\alpha v\beta 3$ receptor ligand, and FB for the α IIb $\beta 3$ receptor. The prototypic ligand **40** (IC₅₀ = 0.7×10^{-9} M) is shown in Figure 18. Generally, the γ -lactam derivatives were highly selective for $\alpha v\beta 3$ versus α IIb $\beta 3$.¹²⁰

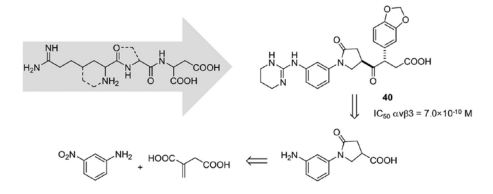


Figure 18 Rationale of the *N*-aryl- γ -lactam strategy and retrosynthesis of potent $\alpha v\beta 3$ antagonist **40**. IC50 value refers isolated integrins.

Molecular-mechanics simulation indicated that the hindered rotation between the *ortho*-substituted central phenyl rings and the lactam imposes to the *N*-aryl- γ -lactam scaffold a well-defined conformation. Relative steric requirements provoke specific arrangements of the guanidine and carboxylate groups, hence influencing the degree of binding to $\alpha\nu\beta3$ integrin. Only the correct conformation of the central core allows the effective interaction with the receptor. The (*R*)-configuration of the γ -lactam was found to be essential for effective binding, as determined by the examination of the individual stereoisomers for other compounds of the series. The favored stereochemistry on the β -amino acid Asp mimic turned out to be the (*S*)-configuration, while chirality modification at the Asp mimic resulted in minor effects on the binding process.

In the last few years, the use of retro sequences of the classic RGD-binding motif for $\alpha\nu\beta3$ and $\alpha5\beta1$ integrin has attracted significant attention.^{40,71,99,101} By contrast, the retro strategy is still largely ignored for the design of ligands of the non RGD-binding $\alpha4\beta1$ integrin. Gentilucci et al.¹⁰² reported the first implementation of the retro-sequence strategy for the design of new analogues of the $\alpha4\beta1$ integrin inhibitor MPUPA-Leu-Asp-Val-Pro-OH **41**, BIO1211,¹²¹ where MPUPA is *o*-methylphenylurea phenylacetyl group (Figure 19). The retro sequences were composed of the β -amino acid (*S*)-pyrrolidine-3-carboxylic acid (or β^2 -Pro) as a constrained core, Asp or *iso*Asp equipped with an aromatic cap at the *N*-terminus, and the amino variant 1-(4(aminomethyl)phenyl)-3-(*o*-methylphenyl)urea (AMPUMP) of the well-known α 4-targeting MPUPA group.¹⁰²

The structures of the novel peptidomimetics were designed on the basis of the models developed by 3D-QSAR analysis of large libraries of compounds containing diphenylureas. This approach led to the retro sequences BnCO-Asp- β^2 -Pro-AMPUMP **43**, and BnCO-*iso*Asp- β^2 -Pro-AMPUMP **44** (Figure 19B). As a variant, retro compounds incorporated an *N*-2,6-dicholorobenzoyl group instead of BnCO. BIO1211 **42** was chosen as $\alpha 4\beta 1$ integrin antagonist reference compound, and the straight sequence MPUPA- β^2 -Pro-AspOBn **42** was examined for comparison purposes (Figure 19A).

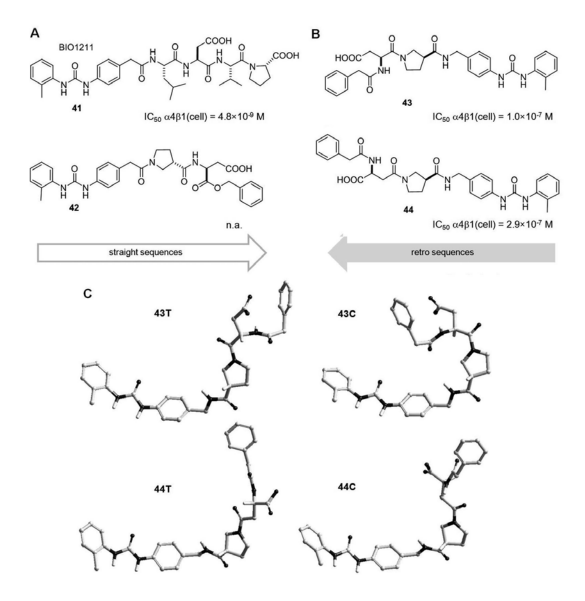


Figure 19 (A) Structures of BIO1211 **41** and the straight sequence containing β 2-Pro **42**. (B) Structures of the retro sequences **43** and **44** containing β 2-Pro. IC50 values refer to integrin-expressing cells (cell). (C) Representative low-energy structure for all-*trans*-**43T** and *trans*-*cis*-*trans*-**43C** (top), all-*trans*-**44T** and *trans*-*cistrans*-**44C** (bottom), consistent with ROESY analysis, and calculated by restrained MD.¹⁰²

The binding of the peptidomimetics to $\alpha 4\beta 1$ integrin was measured by Scintillation Proximity-binding Assay (SPA), using Jurkat cells consistently expressing human $\alpha 4$ integrin, and ¹²⁵I-FN as the specific radioligand. The straight sequence MPUMP- β^2 -Pro-AspOBn **42** was not able to inhibit ¹²⁵I-FN binding. On the contrary, the retro compounds **43**, **44** caused concentration-dependent inhibition of ¹²⁵I-FN binding. The sequence BnCO-Asp- β^2 -Pro-AMPUMP **43** showed IC₅₀ of 1.0×10^{-7} M; the compound **44** containing *iso*Asp showed IC₅₀ of 2.9×10^{-7} M. The analog of **43** equipped with the 2,6-dicholorobenzoyl group revealed modest micromolar affinity.

Subsequently, the ability of the new ligands to inhibit the adhesion of $\alpha 4\beta 1$ integrin-expressing Jurkat cells to VCAM-1 was assayed. The reference compound **41** behaved as a potent $\alpha 4\beta 1$ integrin ligand (IC₅₀ = 7.6×10⁻⁹ M). Interestingly, the retro compounds **43** and **44** inhibited cell adhesion with an IC₅₀ of 80 and 9.3×10⁻⁸ M, respectively, but did not show activity in cell adhesion assays aimed to evaluate potential antagonist activity toward $\alpha \nu \beta 3$ integrin (SK-MEL-24 cells coated with VN). The peptidomimetic **42** and the 2,6dicholorobenzoyl variants of **43** and **44** did not affect cell adhesion to a significant extent, confirming the results of the SPA assay discussed above

The significant biological activity of the structurally related **43** and **44** prompted the analysis of their preferred conformations by NMR spectroscopy in 8:2 DMSO-d₆/H₂O. Cryoprotective mixtures of high viscosity solvents, such as DMSO and water, have been utilized by several authors as excellent biomimetic media for NMR analysis.^{122,123,124} As expected, the ¹H NMR spectra of both compounds revealed two sets of resonances, in about 1:1 ratio, corresponding to the *cis* and *trans* conformers of the peptide bond linking the pyrrolidine ring. Structures consistent with 2D ROESY were analyzed in a box of explicit water molecules by restrained molecular dynamics (MD); the representative geometries of all-*trans*-**43T**, *trans*-*cis*-*trans*-**43C**, all-*trans*-**44T**, *trans*-*cis*-*trans*-**44C**, are shown in Figure 19C.

Despite of the potential high flexibility of the linear ligands, the β -Pro scaffold induced markedly bent conformations. The most significant differences between the predominant conformations of **43** and **44** were found at the position of the aromatic group adjacent to Asp or *iso*Asp, respectively. This might be the consequence of the slightly different affinity values determined by SPA, confirming the relevant role of the flanking aromatic substituent for efficient binding of the receptor.

Apparently, the new compounds are less potent integrin antagonists compared with the reference BIO1211 **41**. However, the latter was previously found to be very unstable in heparinized blood, plasma and rat liver, lung and intestinal homogenates, being metabolized by hydrolytic cleavage of the terminal dipeptide moiety, giving a sequence much less active than the parent compound.^{125,126} Because of their peptidomimetic nature, the sequences BnCO-Asp- β^2 -Pro-urea **43** and BnCO-isoAsp- β^2 -Pro-urea **44** were assumed to be significantly more stable, in particular upon consideration of the presence of the central β -Pro scaffold. The resistance to enzymatic degradation was estimated in mouse serum. After 120 min, **43** was degraded to a moderate extent (about 10%), and **44** was even more stable (<5% degradation) after incubation in the mouse serum. By contrast, BIO1211 **41** was almost completely hydrolyzed under the same conditions.

In this context, Tolomelli et al. employed isoxazolines as the central scaffolds of RGD peptidomimetics. These heterocyclic segments behave as flat, achiral β^2 -proline analogues, which can be derivatized with substituents at the C5 of the pyrrolidine ring. Accordingly, the *trans*-5-hydroxyisoxazolidine-4-carboxylates of interest were prepared via the conjugate addition of hydroxyamine nucleophiles to alkylidene acetoacetates promoted by a Lewis-acid, and subsequent intra-hemiketalization.^{127,128} For instance, racemic *trans*-5-hydroxyisoxazolidine-4-carboxylates were *N*-acylated with benzyl malonyl chloride; mesylation and base-induced elimination gave the corresponding racemic isoxazolines (Figure 20). The coupling of 4-nitrobenzylamine by conventional procedures and the reduction of the nitro group afforded the integrin ligands **45**. Further sequence elongation by introduction of a glycine gave the peptidomimetic **46**.¹²⁹

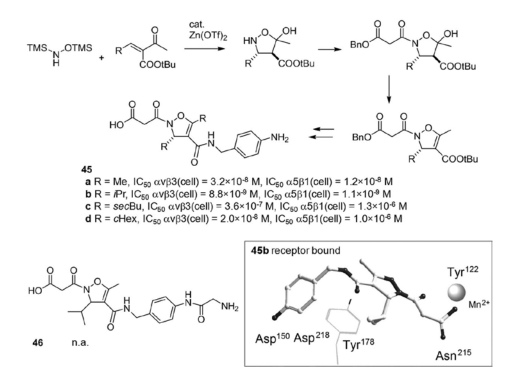


Figure 20 Relevant steps of the synthesis of RGD mimetics **45a-d** and **46** containing the flat isoxazoline core. IC50 values refer to integrin-expressing cells (cell). In the box: sketch of the docking (Glyde version 4.5) best pose of compound (*R*)-**45b** (thick lines) into the crystal structure45 of the extracellular domain of $\alpha\nu\beta3$ integrin (the Mn2+ ion at MIDAS is rendered as a ball). Selected integrin residues involved in interactions with the ligand are highlighted; the stacking interaction between the heterocycle and Tyr178 (rendered in sticks) is also shown.

Experiments were performed to ascertain the ability of racemic compounds **45** and **46** to inhibit the adhesion of $\alpha 5\beta 1$ integrin-expressing cells K562, or the adhesion of $\alpha \nu\beta 3$ integrin-expressing cells SK-MEL-24, to immobilized FN. Among the various RGD mimetics tested, **45a** was an equally potent inhibitor of both $\alpha \nu\beta 3$ and $\alpha 5\beta 1$ integrin, with IC₅₀ in the high nanomolar range; **45b** with R = *i*Pr exhibited the highest inhibitory activity against both $\alpha \nu\beta 3$ and $\alpha 5\beta 1$ integrins with of 8.8 and 1.1×10^{-9} M, respectively. The compound **45c** was poorly active towards both integrin types, while **45d** exhibited selectivity for $\alpha \nu\beta 3$ integrins over $\alpha 5\beta 1$. In contrast, the longer compound **46** was practically inactive. The effect of compound **45b** in integrin signaling *via* ERK was determined. Pre-incubation with **45b** significantly reduced the amount of ERK phosphorylation induced by FN in K562 cells.

A computational model for the binding of **45b** to $\alpha\nu\beta3$ integrin was built by automated docking computations. The ligand was docked into the 3D model of integrin $\alpha\nu\beta3$, as reported in the complex with cilengitide **3** (PDB ID: 1L5G).⁴⁵ Apart from the usual electrostatic interactions anticipated for RGD mimetics

including **3**, the computations highlighted a peculiar stacking effect between the 4-aminobenzylamine moiety of the ligand and the aromatic side chain of Tyr¹⁷⁸ of the α v subunit (Figure 20).

In the course of studies aimed at the development of in-peptide one-step ring closure to give peptidomimetics comprising rigid heterocyclic scaffolds, Gentilucci et al. obtained the 5-aminomethyloxazolidine-2,4-dione (Amo) scaffold.¹³⁰ The Amo ring was obtained in solution or in solid phase from isoserine (*iso*Ser) or other α -hydroxy- β -amino acids already present within the sequence of peptides, by treatment with disuccinimidyldicarbonate (DSC) and a catalytic amount of base (Figure 21A).

This Amo ring incorporated the α -hydroxy- β -amino acid and the amine function of the adjacent residue. This result was complementary to the previously observed reactions with sequences including β -hydroxy- α -amino acids, which gave under the same conditions oxazolidinone (Oxd)-peptides.^{131,132} Indeed, depending on the stereochemistry, linear peptides including two consecutive β -hydroxy- α -amino acids gave rise to sequences Oxd-Oxd or Δ Abu-Oxd (Δ Abu, dehydroaminobutanoic acid). These scaffolds adopted well defined extended or folded conformations, in particular normal or inverse β -turns.¹³³

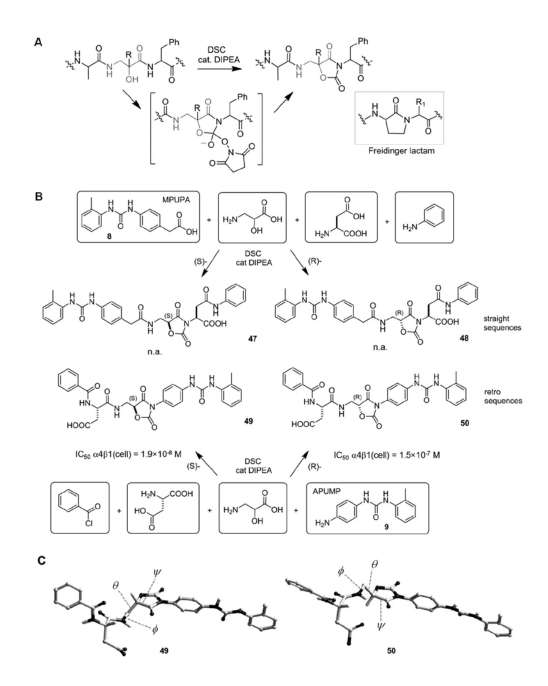


Figure 21 (A) The cyclization of linear peptides containing α -hydroxy- β -amino acids gave Amo dipeptide scaffolds. In the box: a classic Freidinger lactam. (B) Design of the straight Amo peptidomimetics MPUPAAmo-Asp(NHPh)-OH, (*S*,*S*)-**47** and (*R*,*S*)-**48**, and of the retro variants PhCOAsp(OH)-Amo-APUMP, (*S*,*S*)-**49** and (*S*,*R*)-**50**. IC50 values refer to integrin-expressing cells (cell). (C) Representative low-energy structures of **49** and **50** consistent with ROESY analysis and calculated by restrained MD in a box of standard TIP3P water molecules.

To date, the oxazolidine-2,4-dione heterocyclic core has found very little use in medicinal chemistry, the only notable example being a procedure for the synthesis of the 5-aminomethyl-3-oxazolidinone core of the antibiotic linezolid (LNZ) in enantiomerically pure form, *via* an Amo intermediate.¹³⁴ From the structural point of view, the Amo dipeptide represents an unprecedented β^2 -homo analogue of the Freidinger lactam¹¹⁷ (Figure 21A). The convenient synthetic procedure and the peculiar conformational features prompted to utilize the Amo α/β -dipeptide scaffold for the design and synthesis of mimetics of the well-known BIO1211 (**41**)¹²¹ as

 $\alpha 4\beta 1$ integrin inhibitors.¹³⁵ The new peptidomimetics were readily prepared by connecting a diphenylurea pharmacophore and an aryl-substituted Asp, to (*S*)- or (*R*)-configured Amo scaffolds, obtained in turn by the expedient cyclization of the corresponding (*S*)- or (*R*)-isoSer-containing sequences; different assemblies lead to "normal" (**47**, **48**) or retro-sequences (**49**, **50**) (Figure 21B). Only the retro sequences were able to inhibit the adhesion of the $\alpha 4\beta 1$ integrin-expressing Jurkat cells to the ligand VCAM in a dose-dependent manner; in particular the homochiral **49** had IC₅₀ in the 10⁻⁸ M range. Also, the two compounds showed remarkable chemical and enzymatic stability, as determined by incubation in mouse serum.

In the diastereoisomer **49**, the central dihedral angle θ (convention of Banerjee and Balaram⁷) of the Amo residue adopts a +*g* conformation, consistent with the calculated geometries previously reported for β^2 -amino acids. In the diastereoisomer **50** the PhCO-Asp group is positioned at the opposite side of the Amo ring with respect to **49**, and the central dihedral angle θ of Amo adopts a -*g* conformation (due to the opposite stereochemistry, -*g* and +*g* of in (*R*)-Amo and (*S*)-Amo, respectively, correspond). In essence, the Amo scaffold of **49** imposes a preferential conformation, that is compatible with the 3D models (see Section 2) reported in the literature for BIO1211 **41** and similar compounds.

Very recently, BIO1211 **41** mimetics containing a dehydro- β -proline ring have been described.¹³⁶ The scaffolds present in these ligands were synthesized by means of a novel strategy, involving ring closing metathesis of a diallylamino derivative (Figure **22**).¹³⁷ Appendages to the central core were selected on the basis of literature precedent. The synthesized products were assayed for cell adhesion inhibition and in additional assays aimed to ascertain the effect of these purported $\alpha 4\beta 1$ antagonists. A particular ligand, (*R*)-**51a**, was a good integrin ligand, showing IC₅₀ = 1.0×10^{-8} M in the inhibition of $\alpha 4\beta 1$ integrin-expressing Jurkat cells adhesion to VCAM-1. A strong dependence on the stereochemistry of the heterocyclic central core could be observed suggesting a preferred disposition of the lipophilic chain for (*R*) enantiomers. Modest activity was observed toward the homologous integrin $\alpha 4\beta 7$, determined by the inhibition of RPMI8866 cells adhesion to the ECM ligand MadCam-1, while they did not display any activity toward $\alpha 5\beta 1$, $\alpha L\beta 2$, and $\alpha \nu \beta 3$ integrins, analyzed using K-562 cells, Jurkat cells, SK-MEL-24 cells, and the ECM ligands FN, ICAM-1, and FN, respectively.

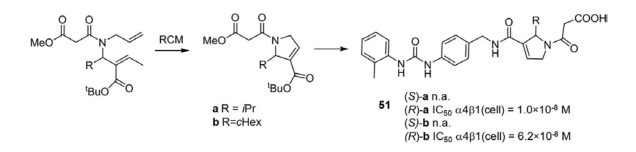


Figure 22 Synthesis of the dehydro- β -proline core, and elaboration of the $\alpha 4\beta 1$ integrin ligands **51**. IC50 values refer to integrin-expressing cells (cell).

5.2.3. Six-membered β-amino acid-derived heterocyclic cores

Some years ago, Singh et al. developed and patented a 3D pharmacophore model of compounds presenting $\alpha 4\beta 1$ inhibitory activity, which was utilized for the design of peptidomimetic ligands.¹³⁸ Among the most promising structures obtained *in silico*, a representative derivative is compound **52** containing the homo- β^2 -proline (nipecotic acid) scaffold (Figure 23).

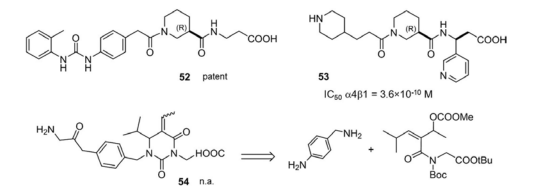


Figure 23 Integrin ligands 52–54 containing 6-membered heterocyclic scaffolds. IC50 value refers to isolate integrins.

The nipecotic acid scaffold is also present in the potent antiplatelet FB receptor (α IIb β integrin) antagonist **53** developed by Hoekstra et al. (Figure 21). Indeed, the six-membered ring scaffold conferred the peptidomimetic ligand an IC₅₀ value for α 4 β 1/FB of 3.6×10⁻¹⁰ M, excellent *in vivo* stability, bioavailability, and adsorption. Oral administration (1 or 3 mg/kg in dogs) produced long-lasting (5-6 h) inhibition of platelet aggregation, but not in the reduction of platelet number.¹³⁹

Finally, Tolomelli et al. synthesized the RGD mimetic **54** containing a dihydropyrimidine-2,4-dione scaffold, formally enclosing a 2-dehydro- $\beta^{2,3}$ -aminoacid and a glycine residue.¹¹³ This compound showed very

poor affinity (IC₅₀ >10⁻⁵ M) for integrins $\alpha v\beta 3$ or $\alpha 5\beta 1$ compared to the structurally related acyclic peptidomimetics.

6. Conclusions

In summary, in this review we present an overview of the state-of-the-art in the design and synthesis of peptidomimetic integrin ligands based on α/β -hybrid peptide structures. While α/β -peptides have been extensively investigated in the foldamer field, scarce attention has been paid so far to these small-molecules from the bioactivity point of view.⁵ Generally, the integrin inhibitors described herein were readily assembled by connecting the pharmacophores to non-racemic β -residues, or to cyclic scaffolds of β -amino acid derivation, usually by expedient procedures.

The different assemblies presented here lead to a variety of 3D diverse structures, and some of the resulting compounds attained remarkable biological activity. The relatively variable receptor affinities were rationalized on the basis of the different 3D molecular geometries imposed by the scaffolds. This evidence supports the fact that the hybrid α/β -peptide approach can furnish bioactive peptidomimetics, potentially useful as diagnostic tools and as therapeutic agents for integrin-correlated diseases.

7. Acknowledgements

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Figure captions

Figure 1. Structures of α -, β^2 -, β^3 -, and $\beta^{2,3}$ -amino acids (following Seebach's nomenclature⁶); the grey arrow shows the extra torsional angle θ (convention of Banerjee and Balaram⁷).

Figure 2. A The RGD and LDVP minimal binding motifs. **B** Structure and extended conformation in solution of **1**. **C** Structure and kinked conformation in solution of **2**. IC_{50} values refer to isolated integrins. The 3D structures were determined by two-dimensional NMR in combination with restrained and free molecular dynamics simulations.³⁷

Figure 3 Structure c[RGDf-*N*(Me)V], cilengitide **3**; in the box: sketch of the main interactions of **3** with the $\alpha\nu\beta$ 3 binding site, as determined from the crystal structure of the complex (PDB ID: 1L5G); the ball represents a Mn⁺² ion.⁴⁵ Di-*N*-methylated **4**, depsi-analogue of cilengitide **5**, and *N*-oligoethylene glycol substituted analogue **6**. IC₅₀ values refer to isolated integrins or to integrin-expressing cells (cell).

Figure 4. Cyclopentapeptides 7 and 8. In the box: the cyclotetrapeptide $\alpha\nu\beta$ 3 antagonist 9, rationally designed by incorporation of a β -amino acid as a *pseudo* γ -turn inducer; the RGD sequence forms a complementary γ -turn; 3D structure of 9 determined by 2D NMR, distance geometry, and restrained molecular dynamics simulations.⁶⁰ Cyclohexapeptide α 5 β 1 antagonist 10. IC₅₀ values refer to isolated integrins or to integrin-expressing cells (cell).

Figure 5. A Structure and backbone conformation of $c[\beta$ -Phe- ψ (NHCO)Ala- ψ (NHCO)Gly-Phe] 11.⁶⁴ B Structure and conformation of $c[(S)-\beta^2$ -Phe-D-Pro-Lys-Phe] 12.⁶³ Conformations determined by 2D NMR and molecular dynamics; some side chains have been omitted for clarity. C Structures of PR-CTP RGD mimetics 13, 14. IC₅₀ values refer to integrin-expressing cells (cell).

Figure 6. Schematic Dunitz-Waser topographic depiction of the eight PR-CTP models (enantiomeric pairs), showing the *S/R* stereochemistry of the C α , and the distances (Å) between the C β atoms. Grey arrows indicate the *pseudo*-axial or *pseudo*-equatorial disposition of the side chains (C α -C β vectors).

Figure 7. Structures of the PR-CTP RGD mimetics **15** and **16**; IC₅₀ values refer to isolated integrins or to integrin-expressing cells (cell). **A**,**B** The α 5 β 1/ α v β 3 mixed integrin antagonist **15** inhibits bFGF-induced formation of capillary-like tubes of HUVEC on Matrigel; **A** HUVEC plated on Matrigel in the presence of

bFGF; **B** HUVEC treated with compound **15** (1 μ M), black bar = 100 μ m. **C** Mean numbers of tubes per square millimeter ± SEM; * P<0.01, ** P < 0.05 vs control (Newman-Keuls test after ANOVA). **D** Overlap of the backbone structures of **15** (black) and **16** (grey), determined by ROESY and molecular dynamics.⁷¹

Figure 8. Sketches of the structures of **15** and **16**⁷¹ determined by molecular docking (Glyde version 4.5) of the ligands within the crystal structure of the integrin $\alpha v\beta 3$ (PDB ID: 1L5G).⁴⁵

Figure 9 A Bioactive conformation of the $\alpha\nu\beta$ 3-selective c[RGDf(*N*-Me)V], cilengitide **3**, showing the distances (Å) between the pharmacophores (grey digits and grey dashed lines) and between C β atoms of Asp, D-Phe, and Arg (black digits and black dashed lines). **B** Dunitz-Waser sketch of **3**. **C** Dunitz-Waser sketches of (*S*,*R*,*S*)-CTP (top), and (*S*,*S*,*S*)-CTP (bottom), models of **15** and **16**, respectively.

Figure 10. Structures of the (+)- and (-)- β -Acc, and β -Acc-containing peptides **17** and **18.** Lowestenergy conformations obtained by restrained MD calculations based on experimental C-C bond distance information. IC₅₀ values refer to integrin-expressing cells (cell).

Figure 11. In the box: structures of cyclic RGD-peptidomimetics 19 and 20 containing diketopiperazine scaffolds, and 3D structure of 20 as obtained by restrained Monte Carlo/Stochastic Dynamics (MC/SD) simulations based on experimental bond distance information, showing a pseudo β -turn at DKP-Arg. Diastereomers 21A and 21B, resulting from hindered rotation. Structure of the c[DKP-RGD]-paclitaxel conjugate 22. IC₅₀ values refer to isolated integrins.

Figure 12. Structure of an integrin antagonist-conjugated MoAb **23**, of a lipid-conjugated integrin antagonist **24**, and of a NODAGA-conjugated integrin antagonist **25**.

Figure 13. Structures of cyclic pentapeptides 26, 27 containing the *iso*DGR motif, and of the cyclic *iso*DGR peptidomimetics 28 and 29 containing the bifunctional DKP scaffolds; 3D molecular structure of 26, 27,^{99,100} and *trans*-(3*S*,6*R*)-DKP-*iso*DGR 29¹⁰¹ obtained by restrained MC/SD simulations based on experimental bond distance information, followed by energy minimization. IC₅₀ values refer to isolated integrins.

Figure 14. Structures of the α IIb β 3 integrin antagonists 30 and 31 containing the trisubstituted β -residues; in the box: retrosynthesis of the β -residues.

Figure 15. General strategies for obtaining sequences 32, 33 containing enantiopure β -amino acids, including the RGD mimetic 34, by CaLB-catalyzed resolution of racemic β -amino acid derivatives. IC₅₀ value refers to integrin-expressing cells (cell).

Figure 16. A Synthesis of enantiomerically pure allylic carbonates and synthesis of dehydro- β -amino esters through amination of allylic carbonates. **B** In the box: ligands **35-37** containing dehydro- β -amino acid scaffolds, examined for integrins $\alpha\nu\beta3$ and/or $\alpha5\beta1$. IC₅₀ values refer to integrin-expressing cells (cell).

Figure 17. Retrosynthetic strategy for the β -lactam-containing RGD-like compounds **38** and **39**. IC₅₀ values refer to integrin-expressing cells (cell).

Figure 18. Rationale of the *N*-aryl- γ -lactam strategy and retrosynthesis of potent $\alpha v\beta 3$ antagonist **40**. IC₅₀ value refers isolated integrins.

Figure 19. A Structures of BIO1211 **41** and of the straight sequence containing β^2 -Pro **42. B** Structures of the retro sequences **43** and **44** containing β^2 -Pro. IC₅₀ values refer to integrin-expressing cells (cell). **C** Representative low-energy structure for all-*trans*-**43T** and *trans*-*cis*-*trans*-**43C** (top), all-*trans*-**44T** and *trans*-*cis*-*trans*-**44C** (bottom), consistent with ROESY analysis, and calculated by restrained MD.¹⁰²

Figure 20. Relevant steps of the synthesis of RGD mimetics **45a-d** and **46** containing the flat isoxazoline core. IC₅₀ values refer to integrin-expressing cells (cell). In the box: sketch of the docking (Glyde version 4.5) best pose of compound (*R*)-**45b** (thick lines) into the crystal structure⁴⁵ of the extracellular domain of $\alpha\nu\beta3$ integrin (the Mn²⁺ ion at MIDAS is rendered as a ball). Selected integrin residues involved in interactions with the ligand are highlighted; the stacking interaction between the heterocycle and Tyr¹⁷⁸ (rendered in sticks) is also shown.

Figure 21. A The cyclization of linear peptides containing α -hydroxy- β -amino acids gave Amo dipeptide scaffolds. In the box: a classic Freidinger lactam. **B** Design of the straight Amo peptidomimetics MPUPA-Amo-Asp(NHPh)-OH, (*S*,*S*)-**47** and (*R*,*S*)-**48**, and of the retro variants PhCOAsp(OH)-Amo-APUMP, (*S*,*S*)-**49** and (*S*,*R*)-**50**. IC₅₀ values refer to integrin-expressing cells (cell). **C** Representative low-energy structures of **49** and **50** consistent with ROESY analysis and calculated by restrained MD in a box of standard TIP3P water molecules.

Figure 22. Synthesis of the dehydro- β -proline core, and elaboration of the $\alpha 4\beta 1$ integrin ligands **51**. IC₅₀ values refer to integrin-expressing cells (cell).

Figure 23. Integrin ligands 52-54 containing 6-membered heterocyclic scaffolds. IC_{50} value refers to isolate integrins.

Biosketches

Rossella De Marco, obtained her *laurea* in Chimistry and Pharmaceutical Sciences (CTF) at the University of Bologna in 2005, and the Ph. D. in Chemical Sciences in 2011 (Synthesis of Modified Amino Acids and Insertion in Peptides and Mimetics. Structural Aspects and Impact on Biological Activity). In 2013 she entered the Young Investigator Programme, Fondazione Umberto Veronesi. Presently, she is interested in the development of new methods for the synthesis of peptidomimetics.

Alessandra Tolomelli graduated in Chemistry in 1994 and received her Ph.D. degree in Chemistry in 1998 at the University of Bologna, working under the supervision of Prof. G. Cardillo. During the Ph.D. she spent a period at the University of California Santa Cruz, working in the group of Prof. J. Konopelsky. She is currently research associate at the University of Bologna. Her research interest mainly focus on beta-amino acids and in their introduction into bioactive peptidomimetics and in their conjugation with diagnostics and cytotoxics.

Eusebio Juaristi was born in Querétaro (México) in 1950 and studied Chemistry at Instituto Tecnológico de Monterrey (B. Sc.) and at the University of North Carolina in Chapel Hill (Ph. D.). Following postdoctoral stays in the University of California-Berkeley and Syntex, California, he returned to Mexico in 1979, to the Chemistry Department of CINVESTAV-IPN, where he is currently Emeritus Professor of Chemistry. He has been Visiting Professor at the ETH-Zurich (1985-1986 and 1992-1993), UC-Berkeley (1999-2000) and RWTH-Aachen (2013). In 1998 he received Mexico's Presidential Medal of Science and in 2006 he became a member of El Colegio Nacional, highest academic distinction in Mexico.

Luca Gentilucci is Associate Professor in organic and bioorganic chemistry at the University of Bologna. He graduate in Chemistry with a thesis on the chemistry of nitrones, and received his Ph. D. with Prof. Cardillo presenting a thesis on the use of heterocyclic intermediates for the synthesis of non-natural amino acids. He expanded the study of azirines and aziridines in Nijmegen with Prof. Binne Zwanemburg. He went back to Bologna as Research Associate for the Interdept. Biotechnology Research Centre, then he returned to the Dept. of Chemistry "G. Ciamician" as Associate Professor. Presently, his main research interests are in the field of the bioactive peptidomimetics (antibiotics, opioids, integrin inhibitors. etc).