Synthesis and Systematic Study on the Effect of Different PEG Units on Stability of PEGylated, Integrin-αvβ6-Specific A20FMDV2 Analogues in Rat Serum and Human Plasma

Kuo-yuan Hung 1, Renata Kowalczyk 1,*, Ami Desai 2, Margaret A. Brimble 1,3,4,*, John F. Marshall 2,* and Paul W. R. Harris 1,3,4,*

1 The School of Chemical Sciences, University of Auckland, 23 Symonds St, Auckland 1010, New Zealand; hungkuoyuan@gmail.com (K.-y.H.); m.brimble@auckland.ac.nz (M.A.B.)
2 Centre for Tumour Biology, Barts Cancer Institute-Cancer Research UK Centre of Excellence, Queen Mary University of London, Charterhouse Square, London EC1M 6BQ, UK; ami.desai89@hotmail.com
3 Maurice Wilkins Centre for Molecular Biodiscovery, University of Auckland, Private Bag 92019, Auckland 1010, New Zealand
4 The School of Biological Sciences, University of Auckland, 3A Symonds St, Auckland 1010, New Zealand
* Correspondence: r.kowalczyk@auckland.ac.nz (R.K.); j.f.marshall@qmul.ac.uk (J.F.M.); paul.harris@auckland.ac.nz (P.W.R.H.)

Abstract: A20FMDV2 is a 20-mer peptide that exhibits high selectivity and affinity for the tumour-related αvβ6 integrin that can compete with extracellular ligands for the crucial RGD binding site, playing a role as a promising αvβ6-specific inhibitor for anti-cancer therapies. Unfortunately, the clinical value of A20FMDV2 is limited by its poor half-life in blood caused by rapid renal excretion and its reported high susceptibility to serum proteases. The incorporation of poly (ethylene glycol) chains, coined PEGylation, is a well-established approach to improve the pharmacokinetic properties of drug molecules. Here, we report a systematic study on the incorporation of a varying number of ethylene glycol units (1–20) into the A20FMDV2 peptide to establish the effects of PEGylation size on the peptide stability in both rat serum and human plasma. In addition, the effect of acetyl and propionyl PEGylation handles on peptide stability is also described. Selected peptide analogues were assessed for integrin-αvβ6-targeted binding, showing good specificity and activity in vitro. Stability studies in rat serum established that all of the PEGylated peptides displayed good stability, and an A20FMDV2 peptide containing twenty ethylene glycol units (PEG20) was the most stable. Surprisingly, the stability testing in human plasma identified shorter PEGs (PEG2 and PEG5) as more resistant to degradation than longer PEGs, a trend which was also observed with affinity binding to integrin αvβ6.

Keywords: A20FMDV2; αvβ6; integrin; PEG; stability

1. Introduction

The integrin αvβ6 is a member of the integrin family that is expressed at low levels in healthy adult epithelia but is up-regulated during foetal development and tissue remodelling such as wound healing [1,2]. Increasing evidence has shown that integrin αvβ6 also promotes tumour growth and metastatic progression of many types of cancer including oral squamous-cell carcinoma, lung, pancreatic, colon, breast and ovarian [3–8]. Ligand binding to integrin αvβ6 is, in part, achieved by binding to the arginine–glycine–aspartate (RGD) motif [9] on specific extracellular ligands such as fibronectin [10], tenascin [11], and the latency-associated peptide (LAP) that forms part of latent-transforming growth factor beta [12,13]. Studies have revealed that synthetic peptides containing a DLXXL (X = any amino acid) subsequence exhibit enhanced affinity to integrin αvβ6 over RGD extracellular...
ligands and exert minimal effects on other integrins [14,15]. These results present a promising opportunity for the development of αvβ6-specific inhibitors as anti-cancer agents.

A20FMDV2 (H2N-NAVPNLRGDLQVLAQKVAR20-T-OH) is a 20-residue peptide derived from the viral protein of foot-and-mouth disease virus [16–19]. This peptide has been shown to exhibit high selectivity and affinity for integrin αvβ6 and can inhibit the binding of extracellular ligands to this integrin [20]. A20FMDV2 binds to integrin αvβ6 through the RGD motif of the 7RGDLQV13L fragment in which the two leucine residues assist the binding of the peptide to the receptor via hydrophobic interactions [21]. The binding of A20FMDV2 to integrin αvβ6 is further stabilised by an α-helix assembled from 10Leu-17Val and flanking C-terminal amino acid residues, thus contributing to the α-helix stability [16,18].

A20FMDV2 labelled with radioactive indium (111In) can be successfully used as an imaging agent for integrin αvβ6 in pulmonary fibrosis [12] and can also serve as a carrier for anti-cancer drugs to form a drug–peptide conjugate targeting integrin-αvβ6-expressing tumour cells [22]. Moreover, the safety of an A20FMDV2-based radioligand for PET studies for use in humans has been reported by Keat et al., when pre-clinical micro-dosing studies on safety, tolerability and immunogenicity were performed on four healthy human subjects using fluorine-18 (18F)-labelled 4-fluorobenzamine (FB)-A20FMDV2 [23]. This shows great promise for A20FMDV2-based ligands to be used in a clinical environment.

However, the use of unmodified A20FMDV2 as a therapeutic agent is limited by its very low half-life due to rapid renal excretion and high susceptibility to serum proteases [24]. We recently undertook a medicinal chemistry approach to improve the pharmacokinetics of A20FMDV2 by the substitution of individual amino acids and the modification of the N- and C-termini [25], while others have employed a decafluorobiphenyl-cyclisation strategy to afford cyclic A20FMDV2 variants that are more stable that their linear counterparts, whilst retaining αvβ6 integrin binding [26].

Another commonly employed modification to improve the pharmacokinetic properties of drug molecules is the incorporation of poly (ethylene glycol) (PEG) chains, a process known as PEGylation [27,28]. PEGs are non-toxic, water-soluble and FDA-approved biopolymers that can minimise the renal clearance and proteolytic degradation of conjugated molecules [29,30]. Synthetic PEGs of large sizes are usually supplied as polydisperse mixtures, affecting batch to batch repeatability during manufacture [31,32]. There is also some evidence that the conjugation of large-molecular-weight PEGs to proteins and biopharmaceuticals increases immunogenicity [33,34]. Nevertheless, PEGylation is a viable option to enhance the pharmaceutical and pharmacological profile of biologically active compounds, and many successful pharmaceuticals on the market are PEGylated [29,30,35,36].

Discrete PEG units such as 1 and 2 (Figure 1A) are readily available and are equipped with an appropriate functional group to permit site-selective PEGylation on amino acid functional groups such as thiols and amines. However, these reagents react indiscriminately with every reactive functional group present on a peptide or protein. Well-defined PEGylating reagents 3–8 and 9–12, respectively (Figure 1B, n = 1–20), are commercially obtainable, monomeric, and bifunctional, hence they can be condensed with the N-terminal amine or side-chain amine of a synthetic peptide. This allows the site-specific incorporation of PEGs via the formation of a stable amide bond and permits the multiple, controlled introduction of PEG units, thereby providing a homogenous PEGylated product. The previous reports on the PEGylation of A20FMDV2 detail the incorporation of large PEG moieties, namely a PEG28 unit covalently attached to the peptide sequence to separate different chemical motifs such as radiotracers within A20FMDV2 [24,37–40].
Studies on A20FMDV2-based position emission tomography (PET) radiotracers incorporating 4-[18F]-fluorobenzoic acid ([18F]-FBA) and PEGs have resulted in beneficial effects of PEGylation on affinity, selectivity and enhanced stability [24,39]. It has been reported that the number and location of PEG moieties within the A20FMDV2 peptide chain (either C- or N- terminus) have different effects on binding to integrin αvβ6 and peptide metabolic stability [24,39].

One study established that selectivity and affinity to integrin αvβ6 for both [18F]-FBA-PEG28-A20FMDV2 and [18F]-FBA-(PEG28)2-A20FMDV2 is not compromised by the incorporation of PEG moieties to A20FMDV2 [24]. Importantly, it was revealed that pharmacokinetic characteristics, measured by the efficiency in tumour retention and washout from the pancreas, was more favourable for αvβ6-targeted pancreatic tumour detection in vivo in a BxPC-3 mouse model [24] using an analogue incorporating only one PEG28 unit. It was also noted that N-terminal A20FMDV2 peptide PEGylation improved the metabolic stability of the construct whilst C-terminal PEGylation increased tumour targeting [39]. These findings suggest that the size and number of the PEG unit itself as well as the location of it within the peptide sequence are important factors when designing PEGylated analogues of A20FMDV2.

In a subsequent study, in vitro and in vivo evaluation of 4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid (DOTA) or 4,11-bis(carboxymethyl)-1,4,8,11-tetraazabicyclo[6.6.2]hexadecane (CB-TEA2)-modified A20FMDV2 analogues incorporating PEG28 showed high affinity and selectivity to αvβ6 integrin, proving a beneficial effect of PEG28 as a spacer between the Cu-chelating DOTA/CB-TEA2 and the peptide itself; importantly, significantly increased specific binding to cells expressing integrin αvβ6 was noted, compared to the analogue lacking PEG28 [37].

More recent studies by Huynh et al., employed PEG28-A20FMDV2-PEG28, which was N-terminally conjugated to either DOTA or 3,6,9,15-tetra-azabicyclo[9.3.1]pentadeca-1(15),11,13-triene-4,5-(4-isothiocyanatobenzyl)-3,6,9-triacetic acid (PCTA) for subsequent radionlabelling with [64Cu] to be used in the imaging or therapy of cancer [41]. The in vitro and in vivo evaluation of the probes’ affinity to αvβ6 in CaSki and BxPC-3 cells, tumour uptake in CaSki and BxPC-3 tumour-bearing mice, and stability in human serum was performed. Promising results for both of the tested probes were reported; however, further optimisation to improve tumour-to-normal-tissue ratios are needed [41].

In another study by the group, gallium-68 ([68Ga])-DOTA-PEG28-A20FMDV2-PEG28 and albumin binding, A20FMDV2-based peptides for the subsequent radionlabelling with

Figure 1. (A) Examples of common PEGylating reagents. (B) Structure of commercially available PEGs containing an acetyl handle (3–8) or a propionyl handle (9–12); n = number of ethylene glycol units.
lutetium-177 ($^{177}$Lu), namely Evans Blue azo dye (EB)-PEG$_{28}$-A20FMDV2-PEG$_{28}$ and 4-(p-iodophenyl)butyric acid (IBA)-PEG$_{28}$-A20FMDV2-PEG$_{28}$, were tested for the αvβ6-specific cancer imaging and therapy [42]. It was shown that the enrichment of the probe with albumin binders increased the blood circulation time, but at the same time non-desired off-target radioactivity accumulation was also observed. The inhibition of tumour growth in mice was also observed; however, toxicity, especially for the Evans Blue azo-dye-enriched analogue, proved to be problematic. Further optimisation of the probes is anticipated to address safety and reduce non-desired normal tissue uptake [42].

Hausner et al., reported that for an A20FMDV2-based radiotracer incorporating a shorter PEG unit (PEG$_{7}$), good stability in rat serum was observed. Unfortunately, compromised tumour uptake was noted in vivo, although this was attributed to the increased lipophilicity of the analogue that incorporated the [$^{18}$F] prosthetic group via copper-free azide–alkyne [3+2]-dipolar cycloaddition [43–45] using the hydrophobic azadibenzocyclooctyne (ADIBO) synthon ([$^{18}$F]FBA-(CH$_2$)$_3$-ADIBO-N$_3$-PEG$_{7}$-A20FMDV2 construct), and not due to the smaller PEG unit [46].

Importantly, [$^{18}$F]-FBA-PEG$_{28}$-[16-Arg]A20FMDV2-PEG$_{28}$ has been recently used in human imaging studies in patients diagnosed with breast, colon, lung or pancreatic cancer, proving to be a promising agent in the early detection of these malignancies due to the high affinity and selectivity for integrin αvβ6 [47]. Additionally, a significant improvement in overall pharmacokinetics was noted compared to the previously reported analogues, including those lacking PEG units [47].

Taken together, the above results demonstrated a therapeutic potential and an improved half-life of A20FMDV2 when either a PEG$_{28}$ or PEG$_{7}$ was incorporated. It should be noted that Li et al., have highlighted the risks associated with higher-molecular-weight PEGylation (5 kDa to 20 kDa) to potentially elicit an immunological response [33].

Many reports exist detailing the positive effect of PEGylation on A20FMDV2 stability when using either larger PEGs or significantly smaller ones (namely PEG$_{28}$, bis-PEG$_{28}$ and PEG$_{7}$). To our best knowledge, no data have been reported thus far where the PEGylation impact on A20FMDV2 stability was investigated in a systematic manner or when using a variety of PEGs other than those that were already reported. The information gained when such studies are performed would be of high relevance to further expand understanding of the PEGylation pattern on A20FMDV2 stability, which could subsequently improve the therapeutic potential of the already clinically relevant A20FMDV2 peptide.

Prompted by the above results, we describe here a systematic study on the effect of increasing the PEGylation of A20FMDV2 as assessed by the measurement of the stability in both rat serum and human plasma, and changes in the specificity and affinity to integrin αvβ6.

Our aim was to identify the shortest PEG unit that improves the A20FMDV2 half-life without compromising its targeting ability to integrin αvβ6. The effect of two different handles, via which PEG units were incorporated into the peptide, namely an acetyl or a propionyl handle, on A20FMDV2 stability was also studied.

Selected PEGylated A20FMDV2 analogues (PEG$_2$–PEG$_{20}$), were subsequently tagged with D-biotin; additionally, diethylenetriaminepenta-acetic acid (DTPA) was incorporated at the N-terminus to allow radiolabelling with indium-111 ([$^{111}$In]). The stability in both rat and human plasma and the specificity and affinity to A375Pβ6 cells (expressing integrin αvβ6) of these compounds using flow cytometry was then assessed. We established that an A20FMDV2 analogue incorporating only five ethylene glycol units (PEG$_5$) displayed preferential characteristics to the non-PEGylated peptide when considering the combined data on human plasma stability and binding ability to integrin αvβ6.

A schematic representation of all the analogues synthesised and tested during the studies is summarised in Table 1.
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Table 1. Synthesized peptides during the studies: A20FMDV2 (13), DTPA-[l²-Ala to Lys(d-biotin)]A20FMDV2 (24) and PEGylated peptide analogues [N-term]-X1-N-X2-VPNLRGDQLQVLAQLVART-OH (14–23, 25–30).

| Compound | N-term. | X1 | X2 |
|----------|---------|----|----|
| 13 (A20FMDV2) | NH₂ | - | Ala |
| 14 | NH₂ | -(CH₂-CH₂-O)₂-CH₂-CO- | Ala |
| 15 | NH₂ | -(CH₂-CH₂-O)₃-CH₂-CO- | Ala |
| 16 | NH₂ | -(CH₂-CH₂-O)₄-CH₂-CO- | Ala |
| 17 | NH₂ | -(CH₂-CH₂-O)₅-CH₂-CO- | Ala |
| 18 | NH₂ | -(CH₂-CH₂-O)₆-CH₂-CO- | Ala |
| 19 | NH₂ | -(CH₂-CH₂-O)₇-CH₂-CO- | Ala |
| 20 | NH₂ | -(CH₂-CH₂-O)₈-CH₂-CO- | Ala |
| 21 | NH₂ | -(CH₂-CH₂-O)₉-CH₂-CO- | Ala |
| 22 | NH₂ | -(CH₂-CH₂-O)₁₀-CH₂-CO- | Ala |
| 23 | NH₂ | -(CH₂-CH₂-O)₁₁-CH₂-CO- | Ala |
| 24 | DTPA-NH | - | Lys(d-biotin) |
| 25 | DTPA-NH | -(CH₂-CH₂-O)₂-CH₂-CO- | Lys(d-biotin) |
| 26 | DTPA-NH | -(CH₂-CH₂-O)₃-CH₂-CO- | Lys(d-biotin) |
| 27 | DTPA-NH | -(CH₂-CH₂-O)₄-CH₂-CO- | Lys(d-biotin) |
| 28 | DTPA-NH | -(CH₂-CH₂-O)₅-CH₂-CO- | Lys(d-biotin) |
| 29 | DTPA-NH | -(CH₂-CH₂-O)₁₀-CH₂-CO- | Lys(d-biotin) |
| 30 | DTPA-NH | -(CH₂-CH₂-O)₂₀-CH₂-CO- | Lys(d-biotin) |

2. Results and Discussion

2.1. Synthesis and Stability Evaluation of A20FMDV2 (13) and PEGylated A20FMDV2 Analogues (14–23) in Rat Serum

PEGylation could, in theory, be carried out at either the N- or C-terminal region of A20FMDV2, or at the side-chain amino group of the native ¹⁶Lys residue. It has been reported that the modification of ¹⁶Lys may afford a peptide with reduced relative activity, most likely due to the disruption of the α-helical region [25]; additionally, N-terminal PEGylation using PEG₂₅₈₆ was shown to be beneficial in improving A20FMDV2 stability over C-terminal PEGylation [39].

Thus, the focus was to establish the effect of the number of ethylene glycol units introduced at the N-terminus of A20FMDV2 peptide on its stability in rat serum.

Commercially available 9-fluorenylmethoxycarbonyl (Fmoc)-protected building blocks 3–12 were used to amiate the N-terminus of A20FMDV2, generating analogues enriched in increasing numbers of ethylene glycol units (1, 2, 3, 4, 5, 8) via an acetyl spacer 14–19. Analogues incorporating 5, 10, 15, and 20 ethylene glycol units via propionyl handle 20–23 were similarly prepared. The stability of the synthetic PEGylated analogues 14–23 was then probed in rat serum and compared to the native, non-PEGylated control A20FMDV2 (13).

The non-PEGylated control 13 and PEGylated A20FMDV2 peptides 14–23 were assembled at a 0.1 mmol scale using aminomethyl polystyrene resin (0.8 mmol/g) synthesised in house [48,49] and manual Fmoc/tert-butyl solid-phase peptide synthesis (Fmoc/IBu SPPS) conditions as described previously and shown in Scheme 1 [25].

To synthesise PEGylated A20FMDV2 analogues 14–23, the resin-bound peptide 31 was amidated using the suitable PEGylating reagent 3–12, namely Fmoc-NH-(CH₂-CH₂-O)₅-CH₂-COOH (n = 1, 2, 3, 4, 5, 8) or Fmoc-NH-(CH₂-CH₂-O)₁₀-CH₂-COOH (n = 5, 10, 15, 20) under HBTU/ DIPEA activation (30 min, rt) followed by the final Fmoc protecting group removal (20% piperidine) prior to the TFA-mediated release of the peptide from the resin Route B). Peptides 14–23 were recovered in >95% purity following purification by RP-HPLC (see Supplementary Materials).
Scheme 1. Synthesis of native A20FMDV2 (13), PEGylated analogues 14–23, DTPA-tagged A20FMDV2 (24), and DTPA-tagged and PEGylated analogues 25–30. Reagents and conditions: (a) Fmoc-Thr(tBu)-HMPPA, DIC, DMF/CH$_2$Cl$_2$ (1:9, v:v), rt, 3 h; (b) 20% piperidine/DMF (v:v), rt, 1 × 5 min + 1 × 10 min; (c) Fmoc-AA-OH, HBTU, DIPEA, DMF, rt, 30 min; (d) TFA, iPr$_3$SiH, DODT, H$_2$O, rt, 3 h; (e) respective PEGylating reagent (3–12), HBTU, DIPEA, DMF, rt, 30 min; (f) DTPA(OtBu), HBTU, DIPEA, DMF, rt, 30 min; (g) 2% hydrazine hydrate/DMF (v:v), rt, 1 × 5 min + 1 × 10 min; (h) D-biotin, HBTU, DIPEA, NMP, rt, 30 min; (i) 3 MBq Indium $^{111}$InCl/nmol, 1 M ammonium acetate, pH 5.5, rt, 30 min.
The stability assay of the native A20FMDV2 peptide (13) and PEGylated derivatives (14–23) was then carried out in rat serum at 37 °C, following previously published procedures [46], and monitored by RP-HPLC over 48 h.

The native peptide (13) was almost completely degraded after 24 h and was undetected at 48 h. For all the PEGylated peptides 14–19 where the PEGs were attached to A20FMDV2 via an acetyl spacer, proteolytic degradation was slower with >30% remaining intact after 48 h (Figure 2A). However, there was no clear trend in that increasing PEGylation did not necessarily result in an increase in peptide stability. Analogue 19 comprising eight ethylene glycol units (PEG8) proved to be the most stable in this series with more than 70% and 58% of the peptide intact at 24 h and 48 h, respectively.

In the PEGylated series 20–23 where a propionyl group was used to space PEGs from A20FMDV2, an improvement in stability compared to the native 13 was also observed.

![Figure 2](image-url)
(Figure 2B). Interestingly, except for the peptide analogue 21 comprising ten ethylene glycol units (PEG10), an increased number of ethylene glycol units (PEG5, PEG15, PEG20) improved the stability of the A20FMDV2 analogues 20, 22, and 23. The A20FMDV2 analogue 23, containing twenty ethylene glycol units (PEG20) was found to be the most stable across all the PEGylated analogues 14–23 overall, with close to 90% and >70% of the peptide remaining intact after 24 h and 48 h, respectively.

Analogues 18 and 20 comprising the same PEG5 component but different chemical handles, i.e., either acetyl (18) and propionyl (20), exhibited a similar degradation pattern in rat serum over the course of 24 h and 48 h (~60% for 18 and 20 after 24 h and ~40% for 18 and 20 after 48 h) suggesting that the length of the linker has no effect on degradation in rat serum.

2.2. Synthesis and Human Plasma Stability of $^{[111]}$In-DTPA-[Lys(d-biotin)]A20FMDV2 (24) and PEGylated $^{[111]}$In-DTPA-[Lys(d-biotin)]A20FMDV2 Analogues (25–30)

Previous studies on the stability of non-PEGylated A20FMDV2 in plasma suggested degradation in mouse serum [20] (50% remaining at 4 h), which significantly differs from the more relevant, clinically used environment of human plasma [25] (>75% remaining at 24 h). Thus, we established a degradation pattern for selected examples of PEGylated A20FMDV2 analogues in human plasma. In this instance, DTPA was incorporated at the N-terminus given the importance of A20FMDV2 as an imaging diagnostic using PET [50], or a single-photon emission computerized tomography (SPECT) that requires the metal chelating DTPA ligand. It was reported that the presence of DTPA on the A20FMDV2 peptide had no significant effect on the specificity and affinity of the peptide, but showed improved stability in mouse serum of $^{[111]}$In-DTPA-[Lys(d-biotin)]A20FMDV2 (50% intact peptide detected at 4 h) [20].

Additionally, to assess PEGylated A20FMDV2 peptide binding activity to the integrin αvβ6 using flow cytometry (see Section 2.3), d-biotin was incorporated via a side chain of the Nε-amino group of lysine-2, substituting the native alanine residue; this modification has previously been well tolerated [20,25,51].

Fmoc/Bu SPPS of the non-PEGylated control 24 and PEGylated analogues of A20FMDV2 incorporating different numbers of ethylene glycol units (i.e., 2, 5, 10, 15, and 20), which were biotinylated at 2Lys and additionally tagged at the N-terminus with DTPA 25–30, was therefore undertaken. In brief, for the preparation of the non-PEGylated analogue 24, the resin-bound and side-chain-protected H2N-Asn(Trt)-Lys(Dde)-Val-Pro-Asn(Trt)-Leu-Arg(Pbf)-Gly-Asp(OBu)-Leu-Gln(Trt)-Val-Leu-Ala-Gln(Trt)-Lys(Boc)-Val-Ala-Arg(Pbf)-Thr(Bu)-HMPPA-PS peptide 32 was directly reacted with Bu-protected DTPA [52] using HBTU/DIPEA in DMF for 30 min at rt, Route C) [25]. For the synthesis of the PEGylated and DTPA-tagged variants 25–30, an initial acylation of 32 with the corresponding Fmoc-NH-(CH2-CH2-O)n-CH2-CH2-COOH (n = 2, 5, analogues 25 and 26, respectively) or Fmoc-NH-(CH2-CH2-O)n-CH2-CH2-COOH (n = 5, 10, 15, and 20), analogues 27–30, respectively), followed by an Fmoc-protecting-group removal by employing the previously described synthetic protocol for the preparation of 13–23 [25], was required, prior to amidation with Bu-protected DTPA, Route D). This was then followed by Nε-1-(4,4-dimethyl-2,6-dioxocyclohexylidene)ethyl (Dde)-protecting-group removal on 2Lys (2% hydrazine hydrate/DMF, v/v) and subsequent attachment of d-biotin to the liberated free Nε amino group using HBTU/DIPEA in 1-methyl-2-pyrrolidone (NMP) for 30 min at rt [25]. Finally, the TFA-mediated peptide cleavage from the resin with simultaneous protecting-group removal afforded the desired constructs 24–30 in good purity (95% or more) following purification by RP-HPLC (see Supplementary Materials).

The DTPA-tagged peptides 24–30 were radiolabelled using 3 MBq $^{[111]}$In chloride/nmol in 1 M ammonium acetate buffer (pH 5.5) (see Methods for details), affording $^{[111]}$In radiolabelled analogues 24–30, Route E). The radiolabelling efficiency and the purity of the radiolabelled compounds were analysed by RP-HPLC.

The $^{[111]}$In-DTPA-A20FMDV2 control 24 and the corresponding PEGylated analogues $^{[111]}$In-DTPA-NH-(CH2-CH2-O)n-CH2-[Lys(d-biotin)]A20FMDV2 (n = 2, 5, 25 and 26,
respectively), and \([^{111}\text{In}]-\text{DTPA-NH-(CH}_2\text{-CH}_2\text{-O})_n\text{-CH}_2\text{-CH}_2\text{[Lys(t-biotin)]A20FMDV2}
(n = 5, 10, 15, and 20) 27–30, respectively) were incubated in either human plasma
or phosphate-buffered saline (PBS) at 37 °C for 24 h, and the samples were analysed using RP-
HPLC, with the results reported as the percentage of peptide remaining after 24 h (Figure 3).

![Figure 3](image_url)

**Figure 3.** Stability of non-PEGylated and PEGylated \([^{111}\text{In}]-\text{DTPA-[Lys(t-biotin)]A20FMDV2}
analogue (24–30) in human plasma. Data expressed as the amount of intact peptide relative to the input
at time zero.

In accordance with our previous studies [25], the non-PEGylated peptide \([^{111}\text{In}]-\text{labelled A20FMDV2 24}
remained with an average of 73% intact after 24 h in human plasma, which significantly differs from
the previously reported stability of this peptide in mouse serum (50% of the peptide remained intact after 4 h) [20].
These data confirm our initial finding that A20FMDV2 is more stable in human plasma than mouse serum [25].

It has been observed in the past that the rates of enzymatic degradation of some
peptides and their analogues differ depending on the biological media in which the stability
testing was performed [53–58]. Among other factors, the observed differences in the rate
of peptide breakdown are most likely due to the species-specific enzyme composition
acting under different mechanisms and pathways [58]. Due to reactivity differences across
media from human and animal species, it is paramount to use the biological medium
of the greatest clinical importance for peptide evaluation. Similar to our observation for
the A20FMDV2 peptide and the PEGylated analogues, Benuck and Marks [54]
showed that rat serum proved to be more active than human serum in the stability studies
of luteinizing hormone releasing hormone (LH-RH), thyrotropin releasing hormone (TRH)
and somatostatin. Our finding further aligns with initial observations that A20FMDV2 is
more stable in human plasma than mouse serum [20] and suggests human-derived media
should be used for subsequent stability studies.

It was observed that the type of handle joining the PEG unit with A20FMDV2 had
a strong effect on peptide stability when testing in human plasma. This was specifically
noted for the \([^{111}\text{In}]-\text{labelled A20FMDV2 analogues incorporating five ethylene glycol units}
(PEG)_5 26 and 27; after 24 h, the analogue 26 utilising an acetyl handle was 80% intact while
analogue 27 bearing a propionyl group was 65% intact.

Importantly, it was noted that the A20FMDV2 analogues incorporating a smaller
number of ethylene glycol units (i.e., PEG_2 and PEG_5) via acetyl handles (25, 26)
displayed a more beneficial effect on the stability in human plasma (around 80% of intact peptide
detected after 24 h) over the analogues that included five or more ethylene glycol units
(i.e., PEG_{10}, PEG_{15}, PEG_{20}) introduced via a propionyl handle (27–30). Within the series
of PEGylated analogues based on the propionyl handle (27–30), degradation showed a linear
effect as the incorporation of an increased number of ethylene glycol units translated to a higher stability, as analogue 30 (PEG20) was ~80% intact after 24 h.

A comparison of the data obtained for the two different sera showed that the type of a handle (acetyl or propionyl) used to attach the PEG unit had no effect on peptide stability when testing was performed in rat serum (e.g., analogues 18-acetyl and 20-propionyl). The stability of the PEGylated analogues incorporating a larger number of ethylene glycol units (20 to 23, PEG5 to PEG20, all propionyl handles) in rat serum gradually increased with a PEG20 analogue being optimal, but in human plasma, the smaller PEG2 and PEG5 (with an acetyl handle) analogues performed the best.

2.3. Specificity and Affinity Evaluation for A375Ppuroβ6 and A375Ppuro Cells of DTPA-[2Lys(D-biotin)]A20FMDV2 (24) and DTPA PEGylated-[2Lys(D-biotin)]A20FMDV2 Analogues (25–30) Using Flow Cytometry

To assess specificity and affinity of the PEGylated A20FMDV2 analogues 25–30 for integrin αvβ6, we used A375Pβ6 cells, which only express integrin αvβ6, and compared this to the binding to A375Ppuro, which contains equal amounts of four RGD-binding integrins (i.e., α5β1, αvβ3, αvβ5, αvβ8) [59].

The PEGylated A20FMDV2 analogues 25–30 were added at a 1000 nM concentration to A375Ppuro or A375Pβ6 cells and binding was measured using flow cytometry. It was shown that none of the peptides bound to the αvβ6-negative A375Ppuro cells, but all the peptides bound equally well to the αvβ6-expressing A375Pβ6 cells, thus confirming that the PEGylated variants retained their absolute specificity for integrin αvβ6 (Figure 4A).

To assess the affinity of PEGylated A20FMDV2 for integrin αvβ6, concentrations of 0.1 nM, 1 nM and 10 nM were compared to the binding of non-PEGylated DTPA-[2Lys(D-biotin)]A20FMDV2 (24).

Across all the concentrations, no loss of activity equated to a value of 1. At the 0.1 nM and 1 nM concentrations, most of the PEGylated peptides lost some affinity compared with the parent peptide (i.e., had a relative value less than 1), particularly those with fifteen and twenty ethylene glycol units (PEG15 and PEG20, 29 and 30, respectively; Figure 4B). However, at the 10 nM concentration, all the PEGylated analogues 25–30 achieved statistically (Anova) similar activities to the parent peptide, revealing that all the PEGylated peptides had good affinity. These data demonstrate that all the PEGylated A20FMDV2 peptides retained complete specificity for αvβ6, but there was some variation in activity at very low concentrations of the peptide.

Hausner et al. [39] reported changes in the in vitro activity of A20FMDV2 that were dependent on the positioning and number of PEG28 units within the A20FMDV2 peptide sequence. The addition of one PEG28 to the N-terminus, or to both the N- and C-termini, retained more activity in vitro than when adding two PEG28 units in series to the N-terminus. It is known that A20FMDV2 binds to integrin αvβ6 through both the RGD motif of the RGDLOV13L sequence and the two leucine residues [16–18]. This binding of A20FMDV2 to αvβ6 is stabilised by an α-helix that is assembled from a C-terminal peptide motif presenting the two non-adjacent leucines as a hydrophobic binding interface [16–18]. The changes in activity observed by Hausner et al. [39] and ourselves (Figure 4) suggest that determining the positioning and number of PEG units within the parent (A20FMDV2) scaffold peptide is critical to minimise the disruption of the ability of the peptide to assume its active conformer. In the work reported here, all the PEGylated A20FMDV2 peptides lost some activity when compared to the native A20FMDV2, but the longer PEGs PEG15 and PEG20 were less tolerated.
Figure 4. (A) Specificity of PEGylated DTPA-[2-Ala to Lys(D-biotin)]A20FMDV2 analogues (25–30) for integrin αvβ6 by flow cytometry; the binding of the peptide (1000 nM) to αvβ6-negative A375Ppuro (white histogram) versus αvβ6-positive A375Pβ6 cells (black histogram). (B) Relative activity of PEGylated DTPA-[2-Ala to Lys(D-biotin)]A20FMDV2 analogues (25–30) for αvβ6 measured by the cell-bound fraction at various concentrations. Data show the binding relative to the same concentration of non-PEGylated DTPA-NK(D-biotin)VPNLRGDLQVLAQKVART as assessed by flow cytometry. Data show mean of three experiments ± SD.
3. Conclusions

A20FMDV2 is a promising lead peptide for cancer treatment as it exhibits high specificity and affinity for the tumour-related integrin $\alpha v\beta 6$ and can inhibit the activation of this integrin by competing with extracellular ligands for the key binding site in integrin $\alpha v\beta 6$. However, the short half-life of A20FMDV2 in blood, caused by the rapid renal excretion and serum proteolytic degradation, suggests that structural modification is required to overcome these limitations. The clinical potential of the PEGylated A20FMDV2 peptide incorporating larger PEG moieties (bis-PEG$_{28}$) has already been realized [47]. It has been reported in the past that the presence of either bis-PEG$_{28}$, PEG$_{28}$ or PEG$_{7}$ within the A20FMDV2 peptide sequence has a positive effect on the peptide pharmacokinetics [24,37–42,46]. The current studies outlined here fulfil the gap in the knowledge where effects of other PEG moieties (PEG$_{1}$-PEG$_{20}$) on A20FMDV2 stability have been investigated with the aim to identify the shortest ethylene glycol repeat that improves peptide stability. The information gained is of great importance to further advance future directions within the A20FMDV2-based drug-discovery pipeline.

Herein, a systematic study was performed on the effect of a discrete number of ethylene glycol units (PEG$_{1}$-PEG$_{20}$) incorporated in an A20FMDV2 peptide at the N-terminus, as well as the choice of the linker (acetyl and propionyl) to couple the corresponding PEG units to the A20FMDV2 peptide, on the peptide stability in rat serum (14–23) and human plasma (25–30). All the peptides could be successfully accessed using Fmoc/tBu SPPS. All the PEGylated (PEG$_{1}$-PEG$_{20}$) A20FMDV2 peptide analogues were less susceptible to proteolytic degradation than the native A20FMDV2 peptide in rat serum. In human plasma, the stability of the PEGylated A20FMDV2 peptide analogues differed significantly from rat serum. The A20FMDV2 analogues comprising a smaller number of ethylene glycol units (PEG$_{2}$ and PEG$_{5}$) were highly stable in human plasma, while the PEGylated PEG$_{20}$ A20FMDV2 peptide proved the most stable in rat serum. We concluded that stability differences in both media underline the importance of the choice of media to more precisely mimic a natural environment of the candidates with clinical potential. It is now evident that human-derived plasma should be the media of choice.

The effect of the chemical handle joining the PEGs with the A20FMDV2 peptide was more pronounced when the degradation studies were performed in human plasma. A PEG$_{5}$-linked acetyl A20FMDV2 was superior to a PEG$_{5}$ propionyl linkage in human plasma, which was more than 80% intact after 24 h; the corresponding propionyl linked PEG$_{5}$ was approximately 65% intact after the same time period. This effect on peptide stability was not observed when the degradation studies were performed in rat serum.

The above observation was also reflected in the binding of the PEG$_{5}$ylated A20FMDV2 peptide to the integrin $\alpha v\beta 6$, as the use of the acetyl handle (26) showed improved binding at 0.1 nM and 1 nM over its propionyl counterpart (27). Taking all the above data into consideration, the PEG$_{5}$-A20FMDV2 analogue 26 containing an acetyl handle had excellent stability in human plasma and retained selectivity and nanomolar affinity for integrin $\alpha v\beta 6$.

Subsequent studies will be undertaken in the future to evaluate promising analogues on cell binding and internalization as well as tumour and normal tissue uptake, and to compare their therapeutic potential with currently existing bis-PEG$_{28}$ A20FMDV2 analogue. These studies will further assess the prospect of simplified PEG modifications on the clinical potential of A20FMDV2 using PEG units that are shorter than those currently employed.

4. Materials and Methods

4.1. Chemistry

All reagents were purchased as reagent grade and used without further purification. The 9-Fluorenylmethoxycarbonyl (Fmoc)-protected amino acids were purchased from CS Bio (Shanghai, China). Fmoc-amino acids were supplied with the following side-chain protection: Fmoc-Asn(Trt)-OH (Trt = triphenylmethyl), Fmoc-Arg(Pbf)-OH (Pbf = 2,2,4,6,7-pentamethyldihydrobenzofuran-5-sulfonyl), Fmoc-Asp(OtBu)-OH (OtBu = tert-butyloxycarbonyl), Fmoc-Gln(Trt)-OH, Fmoc-Lys(Boc)-OH (Boc = tert-butylxycarbonyl), Fmoc-Thr(tBu)-OH. Fmoc-
Thr(tBu)-3-(4-hydroxymethylphenoxy) propionic acid (HMPPA) was purchased from PolyPeptide (Strasbourg, France). Fmoc-NH-(CH$_2$-CH$_2$-O)$_n$-CH$_2$-COOH ($n = 1$–$5$, and 8) and Fmoc-NH-(CH$_2$-CH$_2$-O)$_n$-CH$_2$-CH$_2$-COOH ($n = 5$, 10, 15, and 20) were purchased either from ChemPep Inc. (Wellington, FL, USA) or Peptides International Inc. (Louisville, KY, USA). Fmoc-Lys(Dde)-OH (Dde = 1-(4,4-dimethyl-2,6-dioxocyclohexylidene)ethyl), O-(benzotriazol-1-yl)-N,N,N',N'-tetramethyluronium hexafluorophosphate (HBTU) and N,N'-diisopropylcarbodiimide (DIC) were sourced from GL Biochem (Shanghai, China). N,N'-Diisopropylethylamine (DIPEA), piperidine, 2% hydrazine hydrate, d-biotin, triisopropylsilane ($i$Pr$_3$SiH), 2,2′-(ethylenedioxy)diethanethiol (DODT), formic acid, diethyl ether (Et$_2$O), caffeine, 1-methyl-2-pyrrolidone (NMP), ammonium acetate, ethylenediaminetetraacetic acid (EDTA), phosphate-buffered saline (PBS), ninhydrin, phenol, and potassium cyanide (KCN) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Trifluoroacetic acid (TFA) was purchased from Halocarbon (River Edge, NJ, USA). Dichloromethane (CH$_2$Cl$_2$), and N,N-dimethylformamide (DMF), were purchased from ECP Ltd. (Auckland, New Zealand). Acetonitrile (MeCN) was purchased from Thermo Fisher Scientific (Waltham, MA, USA). Previously published procedures were used for the synthesis of aminomethyl polystyrene resin (0.8 mmol/g) [48, 49] and diethylenetriaminepentaacetic acid tetra tert-butyl ester (DTPA(O$_t$Bu)) [52]. Indium [$_{111}$In] chloride was purchased from Tyco Healthcare UK Commercial Ltd. (Hampshire, UK).

4.2. General Procedure for Peptide Synthesis, Purification, and Analysis

Peptides 13–30 were assembled using manual Fmoc/tBu SPPS (0.1 mmol) using previously published procedures [25] and starting from aminomethyl polystyrene resin (0.8 mmol/g) synthesised in house. PEGylation was carried out using the following procedure: a solution of Fmoc-NH-(CH$_2$-CH$_2$-O)$_n$-CH$_2$-COOH ($n = 1$–$5$, and 8, 3–8) or Fmoc-NH-(CH$_2$-CH$_2$-O)$_n$-CH$_2$-CH$_2$-COOH ($n = 5$, 10, 15, and 20, 9–12) (5 equiv.), HBTU (4.75 equiv.) and DIPEA (10 equiv.) in DMF (1 mL) was added to the resin. After agitating the mixture at rt for 30 min, the solution was drained, and the resin washed with DMF (3 × 5 mL).

After the last step of synthesis, the peptidyl resin was washed thoroughly with CH$_2$Cl$_2$ and dried under the flow of nitrogen. Resin cleavage with simultaneous removal of the amino acid side-chain-protecting groups was undertaken by incubating the resin in TFA/i$Pr_3$SiH/H$_2$O/DODT (v/v/v/v; 94/1/2.5/2.5) cleavage cocktail (5 mL) for 3 h at room temperature (rt). The crude peptides were precipitated and triturated with cold diethyl ether, isolated (centrifugation), dissolved in 50% MeCN (aq) containing 0.1% TFA and lyophilised, prior to their final purification by reverse-phase high-performance liquid chromatography (RP-HPLC).

Peptide purification was performed using a Waters 600 System with a Waters 2487 dual wavelength absorbance detector using a Waters (Milford, MA, USA) Xterra® Prep MS C18 10 μm; 19 × 300 mm preparative column at a flow rate of 10 mL/min. Gradient systems were adjusted according to the elution and peak profiles obtained from the analytical RP-HPLC chromatograms. Fractions were collected, analysed by either RP-HPLC or ESI-MS, pooled and lyophilised. Isolated peptide yields were calculated based on 0.1 mmol synthesis. Analytical RP-HPLC was performed using Thermo Scientific (Waltham, MA, USA) Dionex Ultimate U3000 system (flow rate of 1 mL/min), and a Waters (Milford, ME, USA) Xterra® C18 5 μm; 4.6 × 150 mm analytical column, using a linear gradient of 5% B to 95% B over 30 min, ca. 3% B per minute. The solvent system used was A (0.1% TFA in H$_2$O) and B (0.1% TFA in MeCN) with detection at 210 nm. Direct infusion ESI-MS were recorded on an Agilent Technologies (Santa Clara, CA, USA) 1120 Compact LC connected to an in-line Hewlett Packard (Palo Alto, CA, USA) 1100MSD spectrometer. Samples were introduced using direct flow injection at 0.2 mL/min into an ESI source in the positive mode. The solvent system used was 0.1% formic acid in H$_2$O and 0.1% formic acid in MeCN (1:1, v/v). Major and significant fragments were quoted in the form x m/z (mass to charge ratio).
Peptide radiolabelling was performed according to previously published procedures [20].

5. Biological Studies

5.1. Cell Lines

A375Ppuro (αvβ6-negative) and A375Pβ6 (αvβ6-positive) cell lines were grown as monolayers in Dulbecco’s Modification of Eagles Medium (DMEM; Gibco, Scotland) supplemented with 10% v/v foetal calf serum (Sigma) at 37 °C in a 100% humidified incubator, 8% v/v CO2. All chemicals were purchased from Sigma Aldrich unless otherwise specified.

5.2. Preparation of Peptide

All peptides were dissolved in 0.1% TFA/water to prepare 1 mM stock solutions, which were aliquoted and stored at −20 °C. Thawed samples were diluted directly into cell growth medium.

5.3. Biological Assessment of PEGylated Peptides

To assess specificity and relative activity of peptides, we incubated 2 × 10^5 A375Ppuro or A375Pβ6 cells on wet ice with 1000, 100, 10, 1, 0.1 and 0 nM of each peptide prepared in DMEM supplemented with 0.1% NaN3 and 0.1% bovine serum albumin (0.1/0.1). After 30 min, the unbound peptides were washed away with 0.1/0.1 by centrifugation (twice for 3 min 120 × g). Mouse anti-biotin antibody (Jackson ImmunoResearch Laboratories Inc., West Grove, PA, USA) was added at 1:100 on ice and again washed after 30 min. Rabbit-anti mouse IgG conjugated to Alexafluor 488 (1:250 final dilution; Molecular Probes, Eugene, OR, USA) was added for an additional 30 min on ice before a final washing and resuspension in 500 µL of 0.1/0.1. Samples were analysed by flow cytometry (FACScan, Becton-Dickinson, Franklin Lakes, NJ, USA). Geometric Mean Fluorescence Intensity (MFI) of 10,000 cells was collected. To assess relative changes in affinity, the above experiment was repeated with at least 3 biological repeats and the results were expressed by comparison with Peptide 24 (DTPA-NK(biotin)VPNLRGLQVLAAKVART), which was run at the same time.

6. Rat Serum Stability

The stability assay of A20FMDV2 (13) and its PEGylated derivatives (14)–(23) was performed according to a published protocol [46]. Each peptide was dissolved in a 0.1 M sodium phosphate buffer solution (2.7 mL) (pH 7.4) containing 25% rat serum to yield a final peptide concentration of 20 µM. The resultant solution was mixed, and nine aliquots (270 µL each) were taken and distributed into 1.5 mL Eppendorf vials. The tubes were shaken at 37 °C for different time periods (t = 0 h, 1 h, 2 h, 4 h, 6 h, 8 h, 12 h, 24 h and 48 h), and a 3-fold excess of cold MeCN (810 µL) was added to the vials after each time point and the mixture was placed at 4 °C for 1 h. Caffeine (90 µL of a 0.0002 mg/mL solution in 0.1% TFA in H2O), which was used as an internal standard, was added to the mixture, and the amount of peptide remaining at each time interval was determined as the percentage of the peptide signal, which is calculated as the ratio of the peptide’s peak area under the curve (AUC) to the caffeine AUC, relative to the signal at t = 0 h.

Human Plasma Stability Assays

Human blood was taken from healthy volunteers in sodium heparin tubes and centrifuged at 2000 × g for 10 min. Plasma (300 µL) was incubated with 7.5 MBq of radiolabelled peptide at 37 °C and 150 µL aliquots were analysed either immediately (t = 0) or after 24 h as follows: samples were treated with an equal volume of ice-cold MeCN, then mixed and centrifuged at 14,000 rpm for 5 min. The supernatants were collected and dried using a
centrifugal evaporator for 10 min to remove MeCN. The residuum was filtered through a 0.22 μm filter and analysed by RP-HPLC. Radiopeptides were also incubated at 37 °C with PBS and analysed without further sample preparation.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/molecules27144331/s1: All characteristic data (RP-HPLC and MS) of the synthesized peptides 13–30.

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