A protein microarray-based *in vitro* transglutaminase assay platform for epitope mapping and peptide discovery

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Abstract

Transglutaminases (TGs) are a family of crosslinking enzymes catalyzing the formation of intra- and inter-molecular glutamine-lysine isopeptide bonds in a calcium dependent manner. Activation of transglutaminases is pathogenically associated with severe human diseases including neurodegenerations, cardiovascular diseases, and autoimmune diseases. Although continuous efforts determining the enzymes' substrate preference have been witnessed, a high-throughput assay platform with the "omic" efficiency is still missing for the global identification of substrate-specific TG modification sites. In this study we report a protein microarray-based in vitro TG assay platform for rapid and large-scale (up to 30000 reactions per chip) determination of the glutamine (Q)-bearing TG modification motifs. With this platform we identified the Q16 in superoxide dismutase 1 and Q109 in alpha-synuclein as the primary modification sites for tissue transglutaminase (TG2), the most ubiquitous member of the enzyme family. Of particular interest, we found a close match between the modification motifs and published vaccine epitope sequences in alpha-synuclein. Our data collectively suggest the glutamine and its follow-up five residues on the C terminal of a protein compose a minimal determinant motif for TG2 modification and the TG2 modification motifs determined by our platform could finally correspond to the substrate's epitope sequences in antigen processing. To screen for site-specific interfering peptides and assist gene editing-based protein engineering, we optimized through onchip amino-acid scanning the TG2 modification motif QQIV in the extracellular matrix protein fibronectin and obtained 14 variants with significantly higher TG2 reactivity that might serve as the competitive inhibitor peptides and 1 with

lower reactivity. We further confirmed this optimization approach with 12-mer peptides, the longest ones that could be synthesized on the chip. Taken together, our synthetic transglutaminase assay platform might be able to deliver a precise epitope blueprint for immunotherapeutic targeting and provide proof-of-concept and directional studies for TG-based peptide discovery and protein design.

Introduction

Transglutaminases (TGs) in mammalian cells are a family of crosslinking enzymes posttranslationally catalyzing the covalent isopeptide bond formation between the y-carboxamide groups of a peptide-bound glutamine residue and a free amine or a peptide-bound lysine in a calcium dependent manner¹⁻⁴. Functionally, TG-mediated isopeptide modification results in the stabilization and aggregation of substrate proteins by facilitating the assembly of resistant to proteolysis^{1, 5}. Activation supramolecular structure transglutaminase and TG-mediated isopeptide modification have been demonstrated to be pathogenically associated with severe human diseases neurodegenerations⁶⁻⁹, cardiovascular diseases 10-13, including autoimmune diseases 12, 14, 15. Accumulation of pathogenic proteins crosslinked by transglutaminase has long been recognized as a hallmark of major disease $(AD)^{16-21}$, neurodegenerative disorders including Alzheimer's Parkinson's disease (PD)^{22, 23}, and Huntington's disease (HD)^{24, 25}. In cardiovascular diseases, transglutaminases contribute to the formation of atherosclerotic plaques by crosslinking extracellular matrix (ECM) proteins²⁶⁻³¹ and to hypertensive disorders by modifying and sensitizing vasopressor receptors^{32, 33}. Transglutaminases are also well-known to participate in autoimmune responses through posttranslational modifications favoring the generation of neoantigens³⁴⁻³⁷. Successful amelioration of key disease features by transglutaminase inhibitors in the pre-clinical animal models of related disorders not only establishes TG as an effective therapeutic target but also potentiates transglutaminase inhibitors as useful drugs for the disease treatment³⁸⁻⁴². However, diagnosis of the related complications in patients

usually accompanies a significant build-up of TG-crosslinked aggregates or plaques, and simply inhibiting the enzymes' activity is not sufficient to clean the build-ups. Given this concern, characterizing substrate-specific crosslinking sites and understanding the sequence preference of transglutaminase substrate would be of great importance in designing immunotherapeutic strategies capable of cleaning the aggregates to revoke the disease progression.

To determine the enzymes' substrate preference and map the modification sites among various protein substrates, continued efforts using phage display⁴³⁻⁴⁵, mass spectrometry⁴⁶⁻⁴⁹, protein arrays^{50, 51}, and bioinformatics tools⁵²⁻⁵⁴ have been witnessed. Although these studies greatly advanced our knowledge regarding the modification patterns of transglutaminases, a high-throughput platform with the efficiency of systems biology is still missing for the global identification of substrate-specific TG modification sites. In this study, by combining the tagged amine donor dansyl-cadaverine-based in vitro TG assay⁵⁵ with a protein microarray⁵⁶ we report a platform for rapid and large-scale (up to 30000 reactions per chip) determination of the glutamine (Q)-bearing TG modification motifs. We tested the platform with peptides from neurodegenerative proteins and mapped the primary modification sites for tissue transglutaminase (TG2), the most ubiquitous member of the enzyme family¹⁻⁴, in superoxide dismutase 1 and alpha-synuclein. Our data indicate the glutamine and its follow-up five residues on its C terminal compose a minimal determinant motif for TG2 modification. To manipulate the TG2 modifications on a certain protein and screen for site-specific interfering peptides, we

employed an onchip amino-acid scanning^{57, 58} method for the optimization of

modification motifs. By scanning the TG2 modification motif QQIV in the

extracellular matrix protein fibronectin and 12-mer Q-bearing peptides, the

longest ones that could be synthesized on the chip, we confirmed the

approach's capability to serve peptide discovery and protein engineering.

Methods

Onchip in vitro TG2 assay

To identify the glutamine residues that can be modified by tissue

transglutaminase on the peptide microchip, the synthesized peptide microchip

was incubated with 100 ug/ml guinea pig liver tissue transglutaminase (Sigma)

and 3 mM dansyl-cadaverine (Sigma) in 1 ml of TBS containing 5 mM Calcium

Chloride and 1 mM DTT at 37 degree for 30 minutes. Afterwards, the peptide

chip was washed at least 3 times with TBS. After washing off tissue

transglutaminase and dansyl-cadaverine molecules bound on the synthesized

peptides, the dansyl-cadaverine conjugated on the chip was tracked by rabbit

anti-dansyl antibody (Invitrogen) followed by Alexa Fluor 594-labeled

anti-rabbit secondary antibody (Invitrogen). Fluorescent microchip figures were

quantified and analyzed with ArrayPro32. Original array figures and data are

available upon request.

Results

Mapping TG2 modification sites in neurodegenerative proteins with

high-throughput in vitro assay platform

To establish a high-throughput assay platform for the rapid and large-scale

identification of TG2 modification sites in disease-related proteins, we synthesized on microchips the glutamine-bearing motifs in superoxide dismutase 1 (SOD1) and alpha-synuclein, the pathogenic proteins in amyotrophic lateral sclerosis (ALS) and Parkinson's disease, respectively.

To characterize the glutamine-bearing TG2 modification motifs on these synthesized peptides, dansyl-cadaverine, a well-established amine donor in transglutaminase reaction, was covalently conjugated to the glutamine residues on the peptide chip by purified TG2 with the help of calcium (Figure 1A). After washing off tissue transglutaminase and dansyl-cadaverine molecules bound on the synthesized peptides, the dansyl-cadaverine conjugated on the chip was probed by anti-dansyl antibody followed by Alexa Fluor 594-labeled anti-rabbit secondary antibody. In this way the level of dansyl-cadaverine incorporation on a certain peptide was measured by the fluorescent intensity.

Each peptide synthesized on the chip has a length of at least 4 amino acids, and its maximum length could be up to 12 mer. And the glutamine residue needs to appear in each position of the peptide once to ensure the thorough coverage of the screening (Figure 1B). In this way the peptide screening may also pattern the substrate sequence optimal for the modification. For example, in human alpha-synuclein the surrounding sequence of the first glutamine is AEKTKQ₂₄GVAEAA (Figure 1B). So the sequences for its 4 mer peptides would be QGVA, KQGV, TKQG, and KTKQ, and those for 5 mer would be QGVAE, KQGVA, TKQGV, KTKQG, and EKTKQ. Up to 11 residues on either N

or C side of the glutamine is covered on the chip. Each peptide is named as protein name_Q residue# in the protein_N terminal residue# in the peptide~C terminal residue# in the peptide (In the peptide position of Q is defined as 0, and the one directly before or after it as -1 or 1). So the 4 mer peptide KQ₂₄GV is named as α-synuclein_Q24_-1~2 (Figure 1B). Therefore, for each glutamine residue in a given protein, the initial number of peptide variants synthesized on the chip will be 4+5+6+7+8+9+10+11+12=72. The corresponding peptides with Q to S swap are also synthesized on the same chip as negative controls.

With this approach we identified Q16 in superoxide dismutase 1 and Q109 in alpha-synuclein as the primary modification sites for TG2 (Figure 2B and 2C). In human SOD1 protein there are 3 glutamine residues including Q16, Q23, and Q154. Compared with Q23 and Q154 counterparts, the Q16 peptides with the 5 mer motif directly following the glutamine residue (QGIINF) showed significantly higher fluorescent intensities (>6000), and the Q to S swap could effectively reduce their fluorescent levels, indicating the Q16 residue is the most probable TG2 modification site in the protein (Figure 2B). Similarly, among the 6 glutamine residues in human alpha-synuclein protein, the Q109 residue with its follow-up 5 mer motif (QEGILE) elicited the strongest dansyl-cadaverine incorporation signal and thereby was identified as the TG2 modification site of the protein (Figure 2C). Aligned data with top hits from SOD1 and alpha-synuclein collectively suggest that the Q and its follow-up 5 residues compose a minimal determinant motif for TG2 modification (Figure 2D), which is further confirmed by the peptides with truncated minimal determinant motifs (not shown).

Optimizing TG2 modification sites with the onchip amino-acid scanning

The small peptide QQIV is a transglutaminase substrate identified in the extracellular matrix protein fibronectin. As an amine acceptor, the peptide has been demonstrated to be an effective competitive inhibitor for the transglutaminase reaction⁵⁹. Given this, we chose it as one of the positive substrate peptides on the peptide chip. On our assayed chip the QQIV peptide showed a fluorescent intensity of ~5000 which is much higher than those (~800) of four-residue negative control peptides without glutamine residues and the chip background (Figure 3A). These results confirmed the efficiency and specificity of the dansyl-cadaverine incorporation in our assay system. To scan for a more preferred TG2 substrate and thereby for a more optimized competitive inhibitor than QQIV, we swapped the isoleucine (I) residue in this small peptide to every other amino acid. Except one variant with reduced transglutaminase reactivity, 14 out of the 19 swapped peptides show significantly higher fluorescent signals and the improvement could be up to 3 folds (Figure 3A), suggesting their better candidacy for competitive inhibitor and substrate of TG2 modification than the original QQIV. However, transglutaminase reactivity of most of these 14 mutants was significantly compromised when the hydrophobic V residue was changed to the hydrophilic G (data not shown). To further test the amino acid-scanning approach with the minimal determinant motif for TG2 modification, we swapped to any other amino acid the first leucine residue in the 12-mer small peptide sequence REQLYLDYNVFS, a known TG2 substrate found in a phage display library. Through pan-amino acid scanning at the L residue we found 6 variants (L to N, S, E, R, V or T) with significantly higher transglutaminase reactivity and 6 (L to G, K, M, W, Y or F) with lower reactivity (Figure 3B). Taken together, our result indicates that the residues within the minimal determinant motif could be reasonable targets for the optimization of modification site and the design of substrate-specific interfering peptides.

Elevating peptide's reactivity with transglutaminase by adding glutamine repeats

Characterized in neurodegenerative complications like the polyglutamine diseases, glutamine repeats elicit excellent TG substrate properties in polypeptides by functioning as efficient amine acceptors, and thereby are considered as a biochemical cause for the pathogenesis. To test this feature on our platform, we randomly added the double glutamine repeat motif QQXX (X stands for any of the 20 amino acids) at the N terminal of the peptide QQIV. Consistent with previous findings, the addition of double glutamine repeat motif resulted in a significant upregulation of the TG2-mediated conjugation of dansyl-cadaverine as the average fluorescent intensity (~10000) of the 400 QQXXQQIV peptides is two folds higher than that (~5000) of the original QQIV peptide (Figure 4A) and 297 out of the 400 QQXXQQIV peptides show a fluorescent intensity of more than 5000 on our assayed chip (Figure 4B). With an average fluorescent intensity of ~20000 (Figure 4A), the top 40 QQXXQQIV peptides could serve as competitive inhibitors to block TG2's modification on the QQIV motif of fibronectin. And among them, the peptides with P at the third residue or I at the fourth show up at most times (Figure 4C), indicating a preferred pattern for the linker region between glutamine repeats.

Discussion

In this study we report a protein microarray-based in vitro TG assay platform for fast and high throughput identification of the glutamine (Q)-bearing TG modification motifs among proteins of interest. With this platform we characterized the Q16 in superoxide dismutase 1 and Q109 in alpha-synuclein as the primary TG2 modification sites. Our data indicate the glutamine and its follow-up five residues make up a minimal determinant motif for TG2 modification. To optimize the site for TG2 modification we used an onchip amino-acid scanning method in which the residues near the glutamine residue are swapped with any other amino acid to manipulate its modification level. Using this approach we scanned the isoleucine residue in TG2 modification motif QQIV of fibronectin and obtained 14 variants with significantly higher TG2 reactivity and 1 with lower reactivity. We further confirmed this approach by scanning the first leucine residue within the minimal determinant motif of the 12-mer peptide REQLYLDYNVFS. Consistent with the findings from polyglutamine diseases, we also showed that the addition of glutamine repeats with optimized linker residues are able to significantly elevate the peptide QQIV's transglutaminase reactivity.

Protein aggregation is a hallmark of neurodegenerative diseases including but not limited to Alzheimer's disease, Huntington's disease, and Parkinson's disease, where TG2-mediated crosslinking of the aggregates has been consistently characterized and considered to play a causal and pathogenic role. Therefore, TG2-mediated crosslinking might be able to serve as a crucial

marker for the aggregated proteins. To map the crosslinking sites for the development of immunotherapies targeting the pathogenic aggregates, we designed a protein microarray-based in vitro TG2 assay platform and confirmed its efficacy by identifying the TG2 modification sites in superoxide dismutase 1 and alpha-synuclein. Interestingly, the TG2 modification motifs in alpha-synuclein closely match the antisera epitope sequences of animals immunized with the full length alpha-synuclein^{60, 61}, implying an endogenous role of TG2 modification in the protein's antigen determination. Indeed, the TG2 crosslinking, which results in ubiquitination prevention⁶² and proteolytic resistance, could be a crucial prerequisite for an antigen peptide's proteasomal^{63, 64} or lysosomal⁶⁵ survival in antigen processing. Thus, the TG2 modification motif determined by our assay platform might directly correspond to the substrate's epitope sequence presented by the MHC. And our platform could be a generalized system applying to most epitope mapping efforts. Amyotrophic lateral sclerosis is associated with SOD1 mutations that may greatly alter the protein's reactivity with transglutaminase and thereby contribute to the protein aggregation. With the synthesized mutant peptides on chip, our platform is also able to determine their unique TG2 modification barcodes for immunotherapeutic targeting. To a greater extent, our synthetic assay platform could be used to map the mutation-related crosslinking signatures for disorders frequently associated with genetic mutations like cancer.

Complementing the immunotherapies designed to clear the crosslinked aggregates or plaques, site-specific interfering peptides are competitive

inhibitors that preventively block TG2 modification at certain sites of proteins of interest. Using an *onchip* amino-acid scanning method, we obtained 14 variants of the classic TG2 modification motif QQIV in ECM protein fibronectin with significantly higher transglutaminase reactivity. Since fibronectin crosslinking is associated with atherosclerotic disorders like myocardial infarction and stroke, and organ fibrosis, the variants could be used as the lead candidates for the development of site-specific interfering peptides treating these complications. With the conjugation of proper fluorescent or radioactive molecules, these peptides could be further developed into diagnostic imaging agents tracing the *in situ* build-up of crosslinked plaques. We are also looking forward to applying the *onchip* amino-acid scanning method to the TG2 modification motifs in SOD1 and alpha-synuclein identified in this study. In this way an *onchip* pipeline of mapping-screening would be established for the best efficiency of synthetic and systems biology.

In this study, through scanning the residue directly following glutamine of the minimal determinant motif in the peptides QQIV and REQLYLDYNVFS, we obtained variants with significantly changed transglutaminase reactivity. To generate peptides with modified reactivity, we introduced into the fibronectin peptide QQIV additional glutamine repeats with optimized adaptor sequence as well. With either of the approaches, we manipulated TG2 modification in peptides of interest at our wills. Our results not only show the crucial role of the minimal determinant motif in the regulation of TG2 modification, but also suggest an applicable outlook for our *onchip* peptide optimization assay platform in gene editing-based protein and cell engineering. Thanks to recent

advances (e.g. CRISPR/Cas9) in gene editing technology, we are able to generate *in vivo* mutant proteins with modified transglutaminase reactivity following the initial *onchip* screening in a rapid and large-scale manner. And we aspire to expand the usage of the peptide microarray-based mapping and optimization platform to studies on other posttranslational modifications including but not limited to phosphorylation⁶⁶, ubiquitination^{67, 68}, nitrosylation^{69, 70}, and many others. With the assistance of proper chemical crosslinking reagents like disuccinimidyl tartrate (DST) and formaldehyde, our platform could be further revised to optimize protein-protein interactions by conducting large-scale protein-peptide binding assays⁷¹. In summary, besides epitope mapping for personalized immunotherapy, our platform of synthetic and chemical biology also provides targeting assistance to gene editing activities. We can foresee that in the near future human will be able to direct the evolution of species by on purpose generate genetically modified proteins *in vivo*.

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Figure Legend

Figure 1. Peptide microarray-based *in vitro* TG2 assay platform for rapid and high-throughput identification of modification Q sites among proteins of interest. A. Assay flow chart for the Q mapping platform (Q: glutamine; CD: cadaverine; Dan: dansyl; Ab: antibody; TG2: tissue transglutaminase). B. Peptide layout strategy and naming rule on the microchip using the first glutamine (Q24) motif in human alpha-synuclein as the example. Each peptide where position of Q is defined as 0 and the one directly before or after it as -1 or 1 is named as *protein name_Q residue# in the entire protein_N terminal residue# in the peptide~C terminal residue# in the peptide*.

Figure 2. Identification of the glutamine-bearing TG2 modification sites in SOD1 and alpha-synuclein with the high-throughput *in vitro* assay platform. A. Representative image of the protein microarray-based *in vitro* TG2 assay with the Q peptides in SOD1 showing that the first Q (Q16) peptides (red box) have a generally stronger reaction signal (red box) than the other two (Q23:yellow box; Q154:green box). Q16 residue in SOD1 (B) and Q109 in alpha-synuclein (C) are identified as the primary TG2 modification sites (p<0.05 vs other residues or mutant controls in B and C; n=triplicates; data=mean ± SEM). The top 3 peptide hits of each Q residue in the TG2 reaction are plotted together with their Q to S mutants. D. Alignment of the top 10 hits of Q16 peptides in SOD1 and Q109 peptides in alpha-synuclein indicate their minimal determinant motifs.

Figure 3. Onchip amino-acid scanning generates peptides with

significantly changed reactivity with TG2. A. Pan-amino acid swapping at the I site of the peptide sequence QQIV generates 14 variants (red bars) with significantly higher TG2 reactivity and 1 lower (dark green) (*P<0.05 versus QQIV; n=triplicates; data=mean ± SEM). B. Amino acid scanning at the L residue directly after the Q site of the TG2 substrate peptide REQLYLDYNVFS also obtained variants with significantly changed TG2 reactivity (*P<0.05 versus REQLYLDYNVFS; n=triplicates; data=mean ± SEM).

Figure 4. Addition of glutamine repeat elevates TG2 reactivity of the fibronectin peptide QQIV. A. Addition of glutamine repeat at the N terminal of fibronectin peptide QQIV results significant the in increase dansyl-cadaverine incorporation as measured by fluorescent intensity on the chip (**P<0.01 versus QQIV; X=any of 20 amino acids; n=triplicates; data=mean ± SEM). B. Majority (297 out of total 400) of the QQXXQQIV variants show a higher level of TG2 reactivity as measured by fluorescent intensity on the chip (the fluorescent intensity of QQIV is ~5000 as indicated). C. Among the top 40 QQXXQQIV peptides with the highest fluorescent intensity, the peptides with a P at the third residue or I fourth appear most.

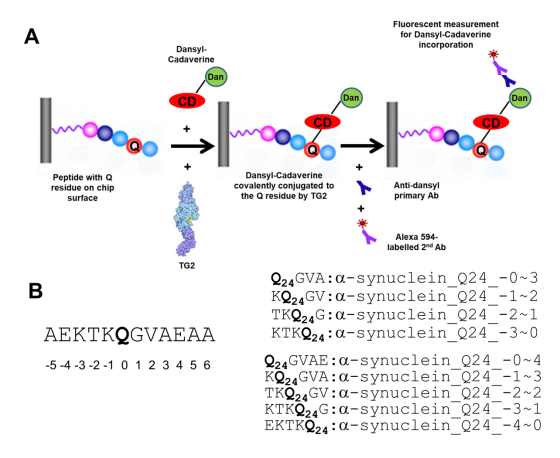


Figure 1

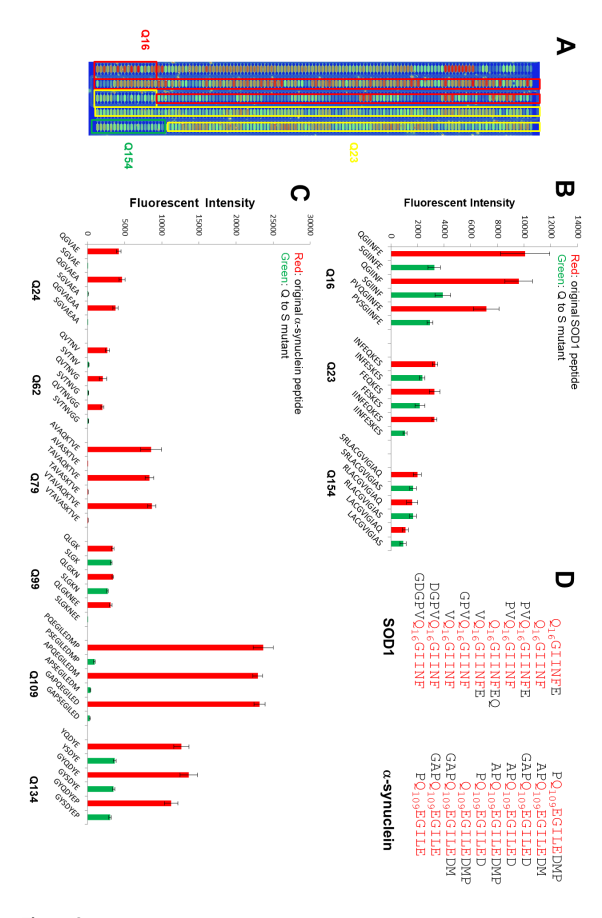


Figure 2

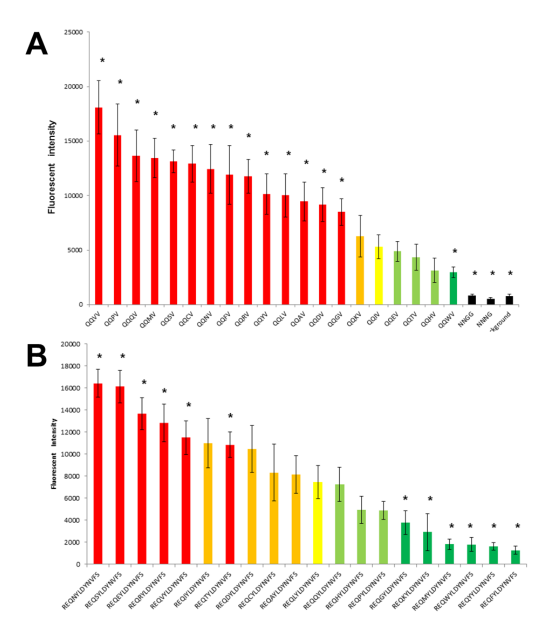


Figure 3

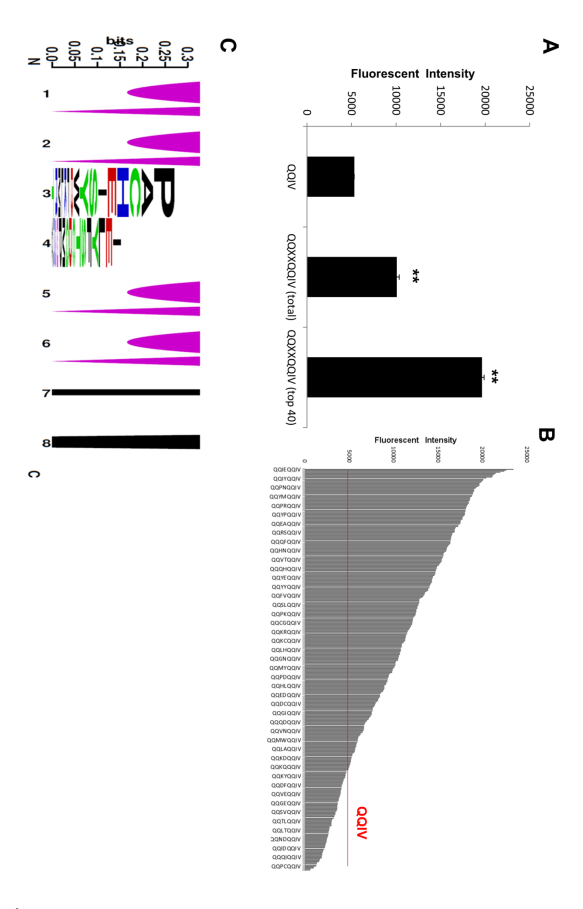


Figure 4