Identification of a single base-pair mutation of TAA (Stop codon) → GAA (Glu) that causes light chain extension in a CHO cell derived IgG1

Taylor Zhang,1,* Yunfu Huang,1 Scott Chamberlain,1 Tony Romeo,1 Judith Zhu-Shimoni,1 Daniel Hewitt,1 Mary Zhu,1 Viswanatham Katta,1 Brad Mauger2 and Yung-Hsiang Kao1

1Protein Analytical Chemistry; Genentech; South San Francisco, CA USA; 2Early Stage Cell Culture Development; Genentech; South San Francisco, CA USA

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Abbreviations: CHO, Chinese hamster ovary; LC-MS/MS, liquid chromatography with tandem mass spectrometry detection; ETS, error-tolerant search; mAb, monoclonal antibody; DTT, dithiothreitol; TFA, trifluoroacetic acid; ESI-TOF, Electrospray-ionization Time-of-Flight; MALDI, Matrix-assisted laser desorption/ionization

We describe here the identification of a stop codon TAA (Stop) → GAA (Glu) = Stop221E mutation on the light chain of a recombinant IgG1 antibody expressed in a Chinese hamster ovary (CHO) cell line. The extended light chain variants, which were caused by translation beyond the mutated stop codon to the next alternative in-frame stop codon, were observed by mass spectra analysis. The abnormal peptide peaks present in tryptic and chymotryptic LC–MS peptide mapping were confirmed by N-terminal sequencing as C-terminal light chain extension peptides. Furthermore, LC-MS/MS of Glu-C peptide mapping confirmed the stop221E mutation, which is consistent with a single base-pair mutation in TAA (stop codon) to GAA (Glu). The light chain variants were approximately 13.6% of wild type light chain as estimated by RP-HPLC analysis. DNA sequencing techniques determined a single base pair stop codon mutation, instead of a stop codon read-through, as the cause of this light chain extension. To our knowledge, the stop codon mutation has not been reported for IgGs expressed in CHO cells. These results demonstrate orthogonal techniques should be implemented to characterize recombinant proteins and select appropriate cell lines for production of therapeutic proteins because modifications could occur at unexpected locations.

Introduction

Sequence variants, defined as protein variants with unintended amino acid incorporation, are potential critical quality attributes for monoclonal antibodies (mAbs) produced for therapeutic use.1 Clone screening to select cell lines free of sequence variants is a general practice to support cell culture development. As a result of the development of more sensitive and systematic methods, several sequence variants have been described.2-4 The process used to detect sequence variants is based on multiple peptide mappings with complementary enzymes, which generates peptides suitable for MS/MS analysis and generally covers 100% of the protein sequence. The peptides are then separated by RP-HPLC coupled with an accurate, high-resolution mass spectrometer. The acquired mass spectra data are subjected to an automated software search, such as Mascot Error Tolerant Search (ETS), against a known protein sequence database. This technique is very sensitive, as sequence variants with serine to asparagine changes due to codon specific mistranslation have been detected at levels as low as 0.1–0.2%.1 Furthermore, the approach can reproducibly detect 0.5% variants as shown by co-mixing two mAbs.2

Sequence variant identification benefits from an array of analytical methods because the protein variants could be detected as abnormal peaks by other methods. Recently, an S441R mutation was identified in another mAb at 0.5% as a result of an investigation of a new peak in the charge assay that was not revealed in the initial ETS.5 In fact, the ETS methodology relies heavily on the correct protein database and might miss a mutation related to stop codons. Even with an ETS performed on a translated DNA database, some spectra are difficult to interpret by software or manually due to limited fragmentation of the peptides. In such a scenario, traditional orthogonal techniques such as N-terminal sequencing will provide invaluable information to identify the variants. Sequence variants can be generated by DNA/RNA mutation or amino acid misincorporation and pinpointing the exact mechanism will enable the design of strategies to avoid sequence variants. Understanding the cause of the variants calls for analyses using nucleic acid based technologies such as PCR or DNA sequencing to complement protein related techniques.6,7

*Correspondence to: Taylor Zhang; Email: yonghuaz@gene.com
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Interestingly, it was reported that 0.2% of mutations among missense and nonsense mutations in the Human Gene Mutation Database (HGMD) are related to stop codons. For example, a point mutation at the stop codon of BRI gene caused a longer open reading frame and generated an extended protein. Protein extension has also been associated with stop codon mutation for apolipoprotein AII (apoAII), which has a 21-residue peptide extension on the carboxyl terminus. Variants with C-terminal extensions for antibodies expressed in CHO cells have not been reported, yet such modifications should be as likely as mutations at any other site. A stop codon-related mutation will generate a C-terminal extension in the protein and needs to be well characterized for biopharmaceutical development because it can affect the function of the expressed proteins.

Here, we report the observation of IgG1 variants with light chain extensions, in addition to the expected 220 amino-acid light chain. The variants were only expressed in one (clone B) of the four clones evaluated during clone screening. With a combination of different enzymatic peptide maps and LC-MS and LC-MS/MS techniques, N-terminal sequencing, RP-HPLC and nucleic acid based technologies, we confirmed that such variants were generated because of a single base-pair mutation of TAA (Stop codon) to GAA (Glu), enabling us to select appropriate clones for clinical therapeutic process development.

Results

Tryptic and chymotryptic peptide mapping revealed extra peptides for clone B. The tryptic peptide mapping chromatographic profiles (Fig. 1) for the four clones are consistent, except there is a new peak eluting at 83 min for clone B. The molecular weight for the new tryptic peptide is 2101.96 Da and it is not an expected tryptic peptide mass according to the antibody sequence. The MS/MS for the peptide had limited daughter ions (data not shown), from which it was difficult to derive the parent peptide sequence. Similarly, the chymotryptic mapping chromatograms (Fig. 2) also revealed a new peak in the clone B antibody digest eluting around 80 min. The mass for the extra chymotryptic peptide is 3546.58 Da, also not an expected chymotryptic peptide mass. Both of the new peaks account for more than 0.1% of the total peak area in their respective maps, indicating the IgG1 produced from clone B has an unknown variant. Mascot search did not provide the identities of these two new peptides. Again manual interpretation of the MS/MS was not successful in identifying the peptide because the peptides are relatively big and fragments present in MS/MS are limited. MS/MS sequencing of these peptides by MALDI yielded no additional information to electro spray ionization MS/MS. Overall, the sequence coverage from these two peptide maps was 100% and there was no other sequence variant found for these four clones.

N-terminal sequencing of the new tryptic and chymotryptic peptides and Glu-C enzymatic peptide mapping confirmed the light chain extension. The two new peptides were fraction collected and subject to Edman N-terminal sequencing. The new peptide collected from the tryptic map was identified as a peptide with the N-terminal as TVAPTECSEAWP (data not shown).
The first eight amino acids are the correct light chain C-terminal sequence indicating protein translation beyond the stop codon. The chymotryptic peptide has the sequence S(C)QV(T)(H)(E)...where parenthesis denotes a tentative sequence call from the N-terminal sequencing data. The results indicate that the extra chymotryptic peptide is also C-terminal peptide (SCQVTHEGSTVEKTVAPTECS) related, though the sequence does not read far enough to show the extension because of the limited sample used. By inspecting the vector sequence near the light chain stop codon, the predicted protein would have the additional sequence XAWPP...continuing until the next in-frame stop codon, 54 nucleotides later. The X here denotes the unknown amino acid resulting from stop codon mutation or mistranslated amino acid. The N-terminal sequence indicates X is glutamic acid, which is consistent with a single nucleotide mutation from stop codon TAA to GAA as shown in Figure 3. The identities of the two new peptides can therefore be deduced as C-terminal light chain peptide extensions. Compared with the correct C-terminal sequence for tryptic peptide TVAPTECS (M+H: 865.36 Da) and chymotryptic peptide SCQVTHEGSTVEKTVAPTECS (M+H: 2308.99 Da) and the new tryptic peak (M+H: 2102.96 Da) and chymotryptic peak (M+H: 3546.58 Da), they have the correct C-terminal light chain sequence plus the same extension of 1237.59 Da, consistent with the calculated mass of the C-terminal extension EAWPPWP...PNLFIAAYNGYK. This peptide eluted at around 91 min because the peptide is more hydrophobic (data not shown).

To further confirm the stop codon mutation to glutamic acid indicated by N-terminal sequencing, we digested protein product from clone B and from a control clone using the enzyme Glu-C. The Glu-C mapping shows a new peak at 30 min for clone B (Fig. 4) compared with the control. The MS data for the related C-terminal peptides are also captured in Table 1. The new peak at 135 min is related to the extension peptide and was marked in the chromatogram. The corresponding MS/MS (Fig. 5) shows the new peptide at 30 min has the y ion series shifted 129 Da compared with the wild type at 29 min; the results are consistent with glutamic acid as the amino acid in the place of stop codon. From the UV peak areas of the wild type peptide KTVAPTECS and mutant peptide KTVAPTECS, we estimate 15% of the C-terminal peptide has the extension, assuming both peptides have the same UV absorptivity.

Knowing that there were light chain extension variants in clone B, we updated the light chain with 18 amino acid extended sequence in the protein database and performed Mascot search again using the peptide mapping data described above. The Mascot software confirmed three peptides with the extended sequence were present in clone B and were absent in the control clone. These peptides are SCQVTHEGSTVEKTVAPTECS EAWPPWP...PNLFIAAYNGYK for the chymotryptic map, KTVAPTECS for the Glu-C map and IAA YNG YK for the tryptic map. The high Mascot score for these peptides (64, 79 and 37 respectively) confirmed these assignments.

Mass spectrometry and RP-HPLC of the mutated variants at the protein level. The mAb was reduced by DTT and analyzed by LC–MS. Deconvolution of the mass data showed the presence of three major light chain species, the expected light chain mass of 23502 Da (1–220 AA, Fig. 6A) and additional masses of 24740 Da and 25493 Da, (Fig. 6B). The 25493 Da species is in agreement with a 17 amino acid extension (1–237 AA) and the 24740 Da species is in agreement with ten amino acid extension (1–230 AA) expected based on translation of the vector DNA sequence with the stop codon mutation to glutamic acid.
The minor mass of the 25622 Da (1–238 AA) species is the result of residual C-terminal lysine on the modified light chain not being completely removed.\textsuperscript{19} The exact cause of light chain ending with phenylalanine (1–230 AA) is unclear, one explanation being CHO cell protease activity.\textsuperscript{20}

Mass analysis of the intact mAbs (Fig. 6C) shows the minor species as 151282 Da compared with the major G0 species of 149289 Da and related glycan variants (149451 G0/G1, 149613 Da (G1/G1 or G2) and the C-terminal lysine variants (149415, 149577 Da). The difference (1993 Da) is in agreement with that of the extension generated by a stop codon mutation to glutamic acid, then translation through to the next in-frame stop codon, assuming the terminal lysine is cleaved from the light chain extension. In Figure 5C, the species of 150,530 Da, which is consistent with intact mAbs with one extended light chain (1–230AA), is also minor.

The IgG1s produced from clone B and control clone were reduced and analyzed by RP-HPLC with UV detection (Fig. 7). Consistent with the peptide mapping results, the new peak in the clone B suggests the mutated light chain is more hydrophobic compared with the normal light chain. The light chain variant peaks comprise about 13.6\% of the total light chain by UV absorption, which is consistent with the results from the Glu-C map.

DNA sequencing confirmed the single base-pair mutation in TAA (stop codon)-GAA (Glu). cDNA generated from clone B and the control clone were submitted to DNA sequencing from both ends of the light chain to allow for reads flanking the base pair in the stop codon (8485 bp) in both directions. The DNA sequencing electrophrogram (Fig. 8A) for clone B showed an additional nucleobase guanine (G) signal that did not surpass the nucleobase thymine (T) signal, but was above baseline noise level, which suggested that a fraction of the mRNA molecules corresponding to the cDNA were mutated from a uracil (U) to a G at the stop codon position in clone B. Thus, the stop codon (TAA) at this location was lost and turned into the codon coding for the amino acid residue glutamic acid (GAA), and protein encoding continued on until the next stop codon. The electropherogram (Fig. 8B) shows that DNA sequence from the control clone has no such mutation. The DNA sequencing results were confirmed with

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure5}
\caption{Tandem mass spectra of (M+2H)\textsuperscript{2+} precursor ions at (A) m/z of 561.75 for peptide KTVAPTECS; (B) m/z of 497.23 for KTVAPTECS. Fragments ions support the conclusion that the light chain stop codon changed to glutamic acid.}
\end{figure}

\begin{figure}[h]
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\includegraphics[width=\textwidth]{figure6}
\caption{Deconvoluted mass spectra of the reduced antibody light chain (A and B) and the intact antibody (C). The theoretical light chain (1–220 AA) is 23502 Da (A) and other light chain species are also observed (B), which is consistent with light chain extension.}
\end{figure}
repeated reactions, and all indicated the mutation in the clone B sequences but not in the control. In addition, this finding is also in agreement with the results from mass spectra data, where a longer coding sequence was obtained from clone B, but not from the control clone.

**Discussion**

With the wide application of accurate and sensitive mass spectrometry, it is possible to detect very small amounts of peptides with alternative sequences down to the 0.01% level—if the sequence variant is known. Nevertheless, peptide mapping with UV detection is still a simple yet powerful method to detect differences in mAbs produced from different clones when the mutation rate is 1 to 5% or above. By UV peptide mapping with complementary enzymes, we detected the extra peaks from the mutated clone in both maps. We could not identify the peaks because of limited fragmentation due to proline-directed fragmentation, but the mutation was at a level much higher than 1%, which allowed us to collect the unknown peaks from tryptic and chymotryptic mapping. The N-terminal sequencing of these collected peptides confirmed the two peptides as the C-terminal light chain extension peptides. Furthermore, the Glu-C map confirmed the stop codon was converted to glutamic acid. The overall approach emphasized N-terminal sequencing as a traditional but critical tool for deciphering the mutation, while the ETS approach was limited to good quality MS/MS data and the correct database. Using ETS, the MS/MS was searched against the custom in-house database generated from ~150 protein sequences. When the mutated peptide sequence is not part of the database, ETS becomes ineffective. After the light chain extension sequence was introduced into the database, Mascot software identified the peptides with good confidence. The finding of a stop codon mutation in our study demonstrates the benefit of establishing the Mascot database with DNA sequence along with the vector sequence. This will help detect a stop codon mutation or N-terminal sequence variants caused by an alternative signal peptidase sequence.

Without DNA sequencing data, it would be impossible to differentiate a stop codon mutation in clone B from a translation error because stop codon read-through could also produce proteins with extensions. It is unclear whether the mutant in clone B originated from a contaminant clone or arose as clone B grew. Theoretically, it is possible that the variant in clone B could result from a transcription error because mRNA was extracted, and cDNA generated and amplified was used for the DNA sequencing; however, transcription errors from DNA occur with low frequency. Because the mutation variants were 15% of the total protein by Glu-C peptide mapping analysis, and the implied nucleic acid mutation by DNA sequencing was at the similar level, it was reasonable to expect that the mutation could be detected and confirmed by Sanger sequencing technology.
which has a well-documented limit of detection (15–20%). Based on the cDNA sequencing data, we selected other clones for further development instead of developing a feeding strategy to eliminate the variants, as demonstrated in another study. To our knowledge, a stop codon mutation on IgG produced by CHO cells has not been reported. This mutation caused an extension of the protein chain. It is also interesting to consider whether codons coding for other amino acids could be mutated to stop codons, which would generate a truncated protein and might be mis-identified as a fragment. In fact, such a mutation has been widely studied in human genetics; incomplete non-functional proteins resulting from such mutations have been associated with the human diseases. We should be cautious about the occurrence of variants resulting from nonsense to sense mutations and focus on the possibility of truncated variants resulting from sense to nonsense mutations because eight triplet codons coding for several amino acids can be converted to the amber stop codon by a single base mutation. These truncated proteins might be removed by Protein A purification; nevertheless, they are statistically more likely than the protein extension observed here. Unlike the typical C-terminal variants such as lysine, proline amidation, or N-terminal signal sequence variants, the light chain mutation variant reported here are even longer, up to 18 additional amino acids. This mutation might increase the risk of immunogenicity because the additional sequence is not part of light chain constant region. Variants with a light chain C-terminal extension might have similar bioactivity to the native protein since the binding is controlled through the variable region in the Fab and the mechanism of action for the intended drug is to bind the antigen, but we did not study the biological activity of the variant. Although clone B had higher titer than the other clones, we eliminated it from further development because of the potential immunogenicity risk. We should note that our observations are valid to stop codons, which would generate a truncated protein and might be mis-identified as a fragment. In fact, such a mutation has been widely studied in human genetics; incomplete non-functional proteins resulting from such mutations have been associated with the human diseases. We should be cautious about the occurrence of variants resulting from nonsense to sense mutations and focus on the possibility of truncated variants resulting from sense to nonsense mutations because eight triplet codons coding for several amino acids can be converted to the amber stop codon by a single base mutation. These truncated proteins might be removed by Protein A purification; nevertheless, they are statistically more likely than the protein extension observed here. Unlike the typical C-terminal variants such as lysine, proline amidation, or N-terminal signal sequence variants, the light chain mutation variant reported here are even longer, up to 18 additional amino acids. This mutation might increase the risk of immunogenicity because the additional sequence is not part of light chain constant region. Variants with a light chain C-terminal extension might have similar bioactivity to the native protein since the binding is controlled through the variable region in the Fab and the mechanism of action for the intended drug is to bind the antigen, but we did not study the biological activity of the variant. Although clone B had higher titer than the other clones, we eliminated it from further development because of the potential immunogenicity risk. We should note that our observations are valid to stable transfection, the DNA encoding the protein of interest is introduced into the genome itself by transfection, and then the clones are subjected to selection and amplification using selective compounds such as methotrexate (MTX), which is a known mutagen. In contrast, transient transfection has the DNA introduced at the beginning of the cell culture run and this DNA is not incorporated into the genome, and the above selection process does not apply.

In conclusion, we identified by multiple analytical techniques an unusual Stop221E mutation in the light chain for an IgG1 produced in CHO cell culture. Our finding emphasizes the importance of orthogonal analytical techniques to support biopharmaceutical development and to characterize unintended modifications in therapeutic proteins.

Materials and Methods

Materials. The IgG1 mAbs were manufactured at Genentech. The antibodies were expressed in four different CHO cell clones listed as A-D and purified by protein A affinity chromatography. Trypsin was purchased from Promega. Chymotrypsin was purchased from Roche Applied Science. Trifluoroacetic acid was purchased from Pierce. HPLC-grade water and acetonitrile were purchased from Burdick and Jackson (Honeywell). Dithiothreitol (DTT) and iodoacetic acid (IAA) were purchased from Sigma.

RCM peptide mapping with Trypsin, Chymotrypsin, Glu-C digestion. Antibody samples were denatured, reduced and alkylated by mixing 50 μL of mAb sample (20 mg/mL) with 10 μL DTT (1 M in Milli-Q water) and 950 μL of the reduction and carboxyamidomethylation (RCM) buffer (6 M guanidine hydrochloride, 360 mM Tris and 2 mM EDTA, pH 8.6), and then mixed with 14 μL IAA (3 M in 1 N NaOH). The reduced and carboxyamidomethylated samples were digested with different enzymes following the typical procedure.

The online LC/ESI-MS/MS analyses of the tryptic digests were performed using an Agilent Capillary 1200 HPLC system coupled with a Thermo Fisher LTQ Orbitrap mass spectrometer equipped with an electrospray ionization source. The antibody digest was injected onto a Waters C4 column with column temperature set at 55°C. The mobile phases and gradient were similar to the published procedure.

N-terminal sequencing. Peptides for N-terminal sequencing were partially dried by vacuum centrifugation. Aliquots were loaded onto Polybrene-polyacryl glass fiber filters and loaded into the reaction cartridge of an Applied Biosystems Procise™ 494HT sequencer. Sequencing was performed using liquid TFA cleavage and PTH-amino acids were identified by comparison with standards using 269 nm absorbance.

Nano chip LC/ESI-TOF MS. Mass analyses of antibodies were performed on an Agilent 6210 electrospray-ionization time-of-flight (ESI-TOF) mass spectrometer (Agilent). Agilent Nano-HPLC Chip (equivalent to Zobax 300SB-C8 analytical column of 43 mm length × 75 μm internal diameter, 5 μm particle size) was used for desalting and separation using a linear gradient from 20% B to 90% B in 6 min. The solvent A was 0.1% formic acid in water and solvent B was 0.1% formic acid in acetonitrile. Spectra consisting of multiple charged protein ions were deconvoluted using the Agilent MassHunter.

RP-HPLC for reduced antibody. RP-HPLC analysis was performed on an Agilent 1200 HPLC system (Santa Clara, CA, USA) equipped with a binary gradient pump, autosampler, temperature-controlled column compartment, and a diode array detector. The system included a Pursuit 3 diphenyl reversed phase column (150 × 4.6 mm, 3 μm, Varian, Lake Forest, CA, USA) that was run at 75°C. The separation was monitored using absorbance at 280 nm. The mobile phase consisted of a gradient of 0.1% TFA in water (mobile phase A) and 0.1% TFA in ACN (mobile phase B). The 46 min method began with a three-minute isocratic hold at 32% mobile phase B, followed by a 26 min linear gradient to 45% mobile phase B. The column was washed at 95% mobile phase B for six minutes and equilibrated at 32% mobile phase B for 10 min.

DNA sequencing to confirm the mutation. The cell cultures were centrifuged in tubes to pellet with a million cells in each pellet and frozen at −70°C. Samples were then processed using the total nucleic acid extraction kit (Qiagen). The RNA was frozen at −70°C until used for RT-PCR by the Onestep RT-PCR kit.
unique results were compiled in Sequencer (Gene Codes) to look for a difference in location flanking the base 5485.

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Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

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