UNITED S	STATES PATENT AND TRADEMARI	X OFFICE
BEFORE	THE PATENT TRIAL AND APPEAL	BOARD
	MYLAN PHARMACEUTICALS INC. Petitioner, v.	
	GENENTECH, INC. Patent Owner.	
	Patent No. 6,407,213	

PETITION FOR INTER PARTES REVIEW

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1001	U.S. Patent No. 6,407,213, Method for making humanized antibodies (filed Jul. 17, 1993) (issued June 18, 2002)
1002 Part I	File History for U.S. Patent No. 6,407,213 Part I
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1004B	Materials Reviewed by Professor Edward Ball, M.D.
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1110	Potter, M. <i>Immunoglobulin-producing tumors and myeloma proteins of mice</i> , Physiol. Rev. 52:631-719 (1972)
1111	Kabat K.A. and Wu, T.T. Attempts to locate complementarity-determining residues in the variable positions of light and heavy chains Ann. NY Acad. Sci. 190:382-93 (1971)
1112	D.R. Davies et al. Antibody-antigen complexes, <i>Ann. Rev. Biochem.</i> 59:439-73 (1990)
1113	Epp et al., The molecular structure of a dimer composed of the variable portions of the Bence Jones protein REI refined at 2.0Å resolution, Biochem. 14:4943 (1975)
1114	Mian, I.S. Structure, function and properties of antibody binding

Exhibit No.	Description
	sites, J. Mol. Biol. 217:133-51 (1991)
1115	Poljak et al. <i>The three-dimensional structure of the fab fragment of a human myeloma immunoglobulin at 2.0-angstrom resolution</i> , Proc. Nat'l Acad. Sci. U.S.A. 71:3440-4 (1974)
1116	Padlan et al. <i>Model building studies of antigen binding sites: The hapten binding site of MOPC315</i> Cold Spring Harbor Symp. Quant. Biol. 41:627-37 (1977))
1117	Boulianne et al. <i>Production of functional chimaeric mouse/human antibody</i> , Nature 312:643-6 (1984)
1118	Padlan, E.A. A possible procedure for reducing the immunogenicity of antibody variable domains while preserving their ligand-binding properties, Mol. Immunol. 28:489-98 (1991)
1119	U.S. Patent No. 6,797,492 Method for Reducing the Immunogenicity of Antibody Variable Domains (veneering of CD18 monoclonal antibodies) (Filed March 16, 2001)(Issued September 28, 2004)
1120	Padlan, Eduardo A., Choosing The Best Framework To Use In The 'Humanization' Of An Antibody by CDR-Grafting: Suggestions From 3-D Structural Data. The 2 nd Annual IBC International Conference on Antibody Engineering. Omni San Diego Hotel, San Diego, CA. (December 16-18, 1991)
1121	Suh et al., <i>The galactan-binding immunoglobulin Fab J539: an X-ray diffraction study at 2.6-Å resolution</i> , Proteins 1:74 (1986)
1122	U.S. Patent No. 5,792,852 <i>Polynucleotides Encoding Modified Antibodies with Human Milk Fat Globule Specificity</i> (humanization of monoclonal antibodies binding to human milk fat globule antigen) (Filed November 16, 1992) (Issued August 11, 1998)
1123	U.S. Patent No. 5,889,157 <i>Humanized B3 Antibody Fragments, Fusion Proteins, and Uses Thereof</i> (humanization of monoclonal antibodies to Lewis [°] -related carbohydrate antigen) (Filed

Exhibit No.	Description
	October 28, 1994) (Issued March 30, 1999)
1124	US Patent No. 5,795,965 Reshaped human antibody to human interleukin-6 receptor (claiming priority to April 25, 1991) (Issued August 18, 1998)
1125	Furey et al. <i>Structure of a novel Bence-Jones protein (Rhe)</i> fragment at 1.6 Å resolution, J. Mol. Biol. 167:661-92 (1983)
1126	Segal et al. <i>The Three-Dimensional Structure of a Phosphorylcholine-Binding Mouse Immunoglobulin Fab and the Nature of the Antigen Binding Site</i> , Proc. Nat'l Acad. Sci. U.S.A. 71:4298 (1974)
1127	Jones, TA Diffraction methods for biological macromolecules. Interactive computer graphics: FRODO, Meth. Enzymol. 115:157-71 (1985)
1128	Co, M. et al. <i>Humanized antibodies for antiviral therapy</i> , Proc. Nat'l Acad. Sci. U.S.A. 88:2869-73 (1991)
1129	History of Microsoft Excel 1978-2013 http://www.exceltrick.com/others/history-of-excel/ (accessed August 29, 2016)
1130	U.S. Patent No. 4,891,762 Method and Apparatus for Tracking, Mapping and Recognition of Spatial Patterns (Filed February 9, 1988) (Issued January 2, 1990)
1131	Wallick, S. et al. Glycosylation of a V_H residue of a monoclonal antibody against $\alpha(1-6)$ dextran increases its affinity for antigen, J. Exp. Med. 168:1099-109 (1988)

I. INTRODUCTION

Pursuant to 35 U.S.C. §§ 311-319 and 37 C.F.R. § 42, Mylan Pharmaceuticals Inc. ("Mylan") petitions for *Inter Partes* Review ("IPR") of claims 1, 2, 4, 12, 25, 29-31, 33, 42, 60, 62-67, 69 and 71-81 of U.S. Patent No. 6,407,213 to Carter, titled "Method for Making Humanized Antibodies" ("the '213 patent," Ex. 1001). With this Petition is a Power of Attorney pursuant to 37 C.F.R. § 42.10(b); and pursuant to 37 C.F.R. § 42.103, the fee set forth in § 42.15(a).

By a preponderance of the evidence, this Petition proves the prior art renders unpatentable claims 1, 2, 4, 12, 25, 29-31, 33, 42, 60, 62-67, 69 and 71-81 of the '213 patent. Methods of making humanized antibodies—the detailed roadmaps in EP0403156 ("Kurrle") [Ex. 1071] and PCT Application No. WO 90/07861 to Queen ("Queen 1990") [Ex. 1050]—anticipate and render obvious to the ordinarily-skilled artisan ("POSITA") the '213 patent's challenged claims as of the priority date. The '213 patent's challenged claims are also obvious in view of Furey [Ex. 1125], Chothia & Lesk [Ex. 1062], Chothia 1985 [Ex. 1063] and/or Hudziak [Ex. 1021].

II. MANDATORY NOTICES

A. Real Parties-In-Interest (37 C.F.R. § 42.8(b)(1))

The real parties-in-interest for Petitioner are Mylan Pharmaceuticals Inc., Mylan Inc., Mylan GmbH, and Biocon Ltd. Mylan N.V. is identified out of an abundance of caution, but this in no way constitutes an admission that it is or was a real party-in-interest in any other IPR proceeding.

B. Related Matters (37 C.F.R. § 42.8(b)(2))

Petitioner is not a party to any litigation related to the '213 patent. The '213 patent is related to the following patents: U.S. Pat. No. 6,639,055; U.S. Pat. No. 6,800,788 (expired, maintenance fee non-payment); U.S. Pat. No. 6,719,971; (expired, maintenance fee non-payment); and U.S. Pat. No. 8,075,890.

C. Identification of Counsel (37 C.F.R. § 42.8(b)(3)) and Service Information (37 C.F.R. § 42.8(b)(4))

Lead Counsel	Back Up Counsel
Jeffrey W. Guise (Reg. No. 34,613)	Deanne M. Mazzochi (Reg. No. 50,158)
Wilson Sonsini Goodrich & Rosati	Rakoczy Molino Mazzochi Siwik LLP
650 Page Mill Road	6 West Hubbard Street, Ste. 500
Palo Alto, CA 94304	Chicago, IL 60654
jguise@wsgr.com	dmazzochi@rmmslegal.com
T: (858)350-2307; Fax: (858)350-2399	T: (312)222-6305; Fax: (312)222-6325

Please direct all correspondence to lead counsel and back-up counsel at the contact information above. Petitioner consents to electronic mail service at jguise@wsgr.com and dmazzochi@rmmslegal.com.

III. GROUNDS FOR STANDING AND PROCEDURAL STATEMENT

As required by 37 C.F.R. § 42.104(a), Petitioner certifies that the '213 patent is available for IPR and that the Petitioner is not barred or estopped from requesting IPR on the grounds identified herein.

IV. IDENTIFICATION OF CHALLENGE AND STATEMENT OF THE PRECISE RELIEF REQUESTED

Petitioner requests *inter partes* review and cancellation of claims 1, 2, 4, 12, 25, 29-31, 33, 42, 60, 62-67, 69 and 71-81 of the '213 patent under pre-AIA 35 U.S.C. §§

102 and 103, as Petitioner's detailed statement of the reasons for the relief requested sets forth. Under 37 C.F.R. § 42.6(c), Petitioner provides exhibit copies, and the Declarations of Dr. Eduardo Padlan (Ex. 1003) and Professor Edward T. Ball (Ex. 1004).

Dr. Eduardo Padlan was a tenure track scientist at the National Institutes of Health, specializing in antibody crystal structure characterization and antibody humanization, which involved identifying key antibody amino acid residues responsible for antigen binding specificity and affinity. His work resulted in numerous publications and patents in this field. Dr. Padlan has also worked as an antibody humanization consultant to many biotechnological companies.

Prof. Edward T. Ball is Professor of Medicine and Director and Chief of the Blood and Marrow Transplantation Division and Program at the University of California, San Diego's School of Medicine. Prof. Ball was an early user of monoclonal antibody therapies, administering mouse monoclonal antibodies to patients beginning in the early 1980s. Prof. Ball has 35+ years of experience in oncology, including developing monoclonal antibody therapies for cancer patients.

The '213 patent's challenged claims generally involve humanized antibodies and humanized antibody variable domains. Ex. 1003 at ¶¶40-63. Claims 1, 2, 4, 12, 25, 29-31, 33, 42, 60, 62-67, 69 and 71-81 of the '213 patent are unpatentable as follows:

Ground	Claims and Basis
1	Claims 1-2, 25, 29, 63, 66, 71, 75-76, 78, 80-81, anticipated by Kurrle
2	Claims 1-2, 4, 29, 62-65, 80-81, anticipated by Queen 1990
3	Claims 1-2, 4, 25, 29, 62-67, 69, 71-72, 75-76, 78, 80-81 as obvious over
	Queen 1990 and Kurrle
4	Claim 12 obvious over Queen 1990 and Kurrle, and also in view of Furey
5	Claims 73, 74, 77 and 79 as obvious over Queen 1990 and Kurrle, and
	also in view of Chothia & Lesk and Chothia 1985
6	Claims 30, 31 and 33 as obvious over Queen 1990, in view of Hudziak
7	Claim 42 as obvious over Queen 1990, in view of Hudziak and Furey
8	Claim 60 as obvious over Queen 1990, in view of Hudziak and Chothia &
	Lesk

V. THRESHOLD REQUIREMENT FOR INTER PARTES REVIEW

A petition for IPR must demonstrate "a reasonable likelihood that the petitioner would prevail with respect to at least 1 of the claims challenged in the petition." 35 U.S.C. § 314(a). As explained below, there is more than a reasonable likelihood that Petitioner will prevail with respect to at least one of the challenged claims.

VI. STATEMENT OF REASONS FOR THE RELIEF REQUESTED

A. Summary of the Argument

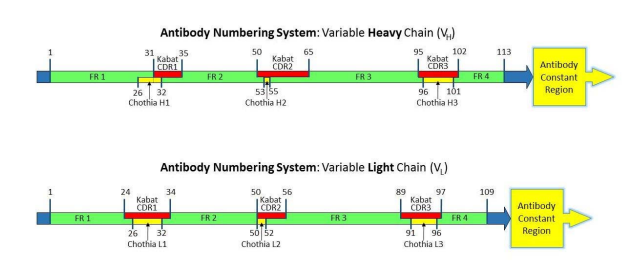
In 1975, the journal *Nature* published Köhler and Milstein's ground-breaking study manufacturing "predefined specific antibodies by means of permanent tissue culture cell lines." Ex. 1022 at 1. Mouse monoclonal antibodies exhibited therapeutic and diagnostic promise, but researchers discovered patients receiving mouse antibodies experienced a human anti-mouse antibody (HAMA) immunogenicity response. Ex. 1003 at ¶70-72; Ex. 1004 at ¶45.

To neutralize the HAMA response, mouse antibodies were first re-engineered

to make them "more human" by replacing parts of the mouse antibody with human counterparts. First generation (early 1980s) versions replaced the mouse antibody's constant region with corresponding human antibody residues. Ex. 1003 at ¶89-91; Ex. 1004 at ¶46-47. While "chimeric" antibodies retained the parent mouse's affinity and specificity, patients still experienced HAMA responses from the mouse variable domain. Next, scientists replaced mouse variable domain framework regions (FR) flanking the complementarity determining regions (CDR) with human sequences. Ex. 1003 at ¶92.

However, because adding human FRs to the regions between the mouse CDRs was known to disrupt binding affinity, the next logical step in the evolution of humanized antibody technology was to switch select residues in the human FRs back to the mouse residue. Ex. 1003 at ¶¶92-107. These techniques were well-known and well mapped out prior to the earliest priority date (June 14, 1991) of the '213 patent. *Id.* at ¶107. Kurrle [Ex. 1071] is just one example disclosing combining human FRs with mouse CDRs, wherein select residues in the human FRs were switched back to mouse. Kurrle's switched residues include claimed residues **4L**, **69H**, **71H**, **73H** and **76H**. Ex. 1071 at 3, Il. 9-10. Kurrle's result was "essentially a human antibody with a much lower immunogenicity in patients." *Id.* at 3, 11-12. Kurrle thus anticipates claims 1, 2, 25, 29, 63, 66, 71, 75, 76, 78, 80 and 81. Ex. 1003 at ¶¶142-158.

Queen 1990 established a humanization roadmap with four specific yet universal criteria for producing humanized antibodies from non-human monoclonal antibodies, including substituting for the mouse monoclonal antibody residue in the Framework Regions (FR) "immediately adjacent to one or more of the 3 CDR's in the primary sequence" according to the Kabat numbering system. Queen 1990 [Ex. 1050]. Kabat [Ex. 1052] and Chothia & Lesk [Ex. 1062] had earlier classified the antibody variable domain structure, including defining the boundaries of the Kabat CDRs, the Chothia hypervariable regions and FRs [Ex. 1003 at ¶¶73-91, 115-16]:



These defined FR/CDR border positions readily allow a POSITA, given Queen 1990's instruction to substitute CDR-adjacent FR residues, to identify at least claimed residues **36H** and **98L** (*see* claims 1, 2, 4, 29, 62, 65, 64, 80 and 81). Queen 1990 thus anticipates at least claims 1, 2, 4, 29, 62, 65, 64, 80 and 81 of the '213 patent under the claims' broadest reasonable construction. Ex. 1003 at ¶¶159-90.

Moreover, all challenged claims—whether e.g., they broadly or more

specifically list residues or properties, or particular antibodies to humanize, are obvious given the prior art, including Queen 1990; Kurrle; and others. Ex. 1003 at ¶¶191-242; 333-43. A number of prior art references and preeminent researchers in the field, including Professors Cyrus Chothia and Arthur Lesk, taught the importance of specific claimed residues, and their predicted contribution to antigen binding, including **93H**, **78H** and **66L**. Ex. 1003 at ¶¶88, 110-20. Their inclusion in the challenged claims thus was not a patentable advance in the field, but obvious.

The prior art also disclosed both p185^{HER2} as a promising therapeutic target, and a specific monoclonal antibody (4D5) against the p185^{HER2} target. Mylan's experts, Professor Ball and Dr. Padlan, both agree that the next logical and necessary step in the development of 4D5 was humanizing it. Ex. 1003 at ¶¶318-23; Ex. 1004 at ¶¶63-79. Queen 1989 and 1990 provided motivation and a sufficient roadmap to accomplish this humanization. Ex. 1003 at ¶¶324-32. Others gave further details on specific residues. *Id.* Thus, given Queen 1990 and Kurrle, or in combination with other references as detailed below, the challenged '213 patent claims also were obvious.

B. '213 Patent-Background

1. The '213 Patent

The '213 patent issued June 18, 2002 from a continuation-in-part of an earlier-abandoned U.S. Patent Appl. No. 07/715,272 (filed June 14, 1991), the '213 patent's

earliest possible priority date.

The '213 patent issued with 82 claims. Claims 1, 30, 62-64, 66, 79 and 80 are independent claims, and all claim a "humanized antibody," "antibody," humanized variant of a non-human parent antibody" or "humanized antibody variable domain" comprising a "non-human ... CDR," and a "Framework Region [FR] amino acid substitution" reverting substituted human framework residues back to, e.g., mouse, at "a site selected from the group consisting of" certain recited residues. Claim 1 chooses from 14 FR light chain residues (4L, 38L, 43L, 44L, 58L, 62L, 65L, 66L, 67L, 68L, 69L, 73L, 85L, 98L); and 10 heavy chain residues (2H, 4H, 36H, 39H, 43H, 45H, 69H, 70H, 74H, and 92H) under Kabat's numbering system. Ex. 1001 at col. 86, Il. 44-52. Claims 30, 62 and 63 add 4 FR residues to claim 1's list (46L, 75H, 76H and 78H). Claim 30's antibody "binds p185^{HER2} and comprises a humanized antibody variable domain"; claim 63's humanized antibody "lacks immunogenicity compared to a non-human parent antibody upon repeated administration to a human patient" when treating chronic disease. Id., col. 88, 11. 27-48.

Claim 66 offers a different list of 5 FR residues: 24H, 73H, 76H, 78H and 93H. *Id.* at col. 88, l. 66 to col. 9, l. 6. Claim 79 lists 4 FR "substitutions at heavy chain positions 71H, 73H, 78H and 93H, utilizing the numbering system set forth in Kabat." *Id.* at col. 90, ll. 3-10. Claim 80 claims the residues of claim 1 plus the 5 residues from claim 66, and adds that the FR amino acid substitution: "(a) noncovalently binds

antigen directly; (b) interacts with a CDR; or (c) participates in the V_L - V_H interface by affecting the proximity or orientation of the V_L and V_H regions with respect to one another." *Id.* at col. 90, Il. 11-25.

Claim 64's "humanized variant of a non-human parent antibody" includes "the most frequently occurring amino acid residues at each location in all human immunoglobulins of a human heavy chain immunoglobulin subgroup wherein amino acid residues forming Complementarity Determining Regions (CDRs) thereof comprise non-human antibody amino acid residues, and further comprises a Framework Region (FR) substitution where the substituted FR residue: (a) noncovalently binds antigen directly; (b) interacts with a CDR; (c) introduces a glycosylation site which affects the antigen binding or affinity of the antibody; or (d) participates in the V_L - V_H interface by affecting the proximity or orientation of the V_L and V_H regions with respect to one another."

The dependent claims recite specific residues (claims 12, 25, 42, 60 and 71-77; claims 75-77 further add a substitution at residue 71H); that the substituted humanized antibody residue is "found at the corresponding location of the non-human antibody from which the non-human CDR amino acid residues are obtained" (claims 2, 31, 67 and 81); that the human antibody variable domain is a "consensus" domain (claims 4, 33 and 69); or an antibody comprising the claimed humanized variable of claims 1 or 66 (claim 29 and claim 78, respectively).

The patent specification's humanization concepts were neither new nor unknown. It acknowledges the widely held view that the "function of the antibody is dependent on its three dimensional structure, and that amino acid substitutions can change the three-dimensional structure of an antibody" near the CDRs. *Id.* at col. 3, ll. 40-44. It acknowledges past "molecular modeling" had "increase[d] the antigen binding affinity of a humanized antibody." *Id.* at col. 3, ll. 44-48. The '213 patent applies the same cloning and analysis tools and techniques that Kurrle [Ex. 1071] and Queen 1990 [Ex. 1050] already described, including site-directed mutagenesis, molecular modeling and antibody functionality analysis. The '213 patent likewise recognizes the known existing promise of p185^{HER2} monoclonal antibody (4D5) as a therapeutic anticancer agent whose murine origin may render it "immunogenic in humans." *Id.* at col. 3, l. 56 to col. 4, l. 23.

2. Brief Overview of the '213 Patent's Prosecution History and Related Proceedings in the PTO

'206 Application Prosecution. The '213 patent issued from Application No. 08/146,206 ("the '206 application"). During prosecution, the PTO rejected the '206 application's claims for anticipation, obviousness, lack of written description, lack of enablement, indefiniteness and non-statutory obviousness-type double patenting. The PTO's unpatentability bases included Queen 1989 [Ex. 1034] and Kabat 1987 [Ex. 1052], asserted here.

Interference with Application No. 11/284,261. Applicants for Application No. 11/284,261 ("the '261 application" or "Adair") requested an interference with the '213 patent. The interference count was drawn to humanized antibodies with non-human substitutions at specific variable domain framework positions. The Board declared the interference, identifying the claims corresponding to the count as claims 30, 31, 60, 62, 63, 66, 67, 70, 73, 77-81 of the '213 patent and claim 24 of Adair. Carter v. Adair, Interference No. 105,744, Declaration of Interference at 4 (Feb. 2, 2010) [Ex. 1095].

The Board determined, however, that claims in an earlier application, to which Adair claimed priority, did not provide pre-critical date support for claim 24, thereby concluding that Adair's claim 24 was barred under 35 U.S.C. § 135(b)(1). Decision on Motions at 9-10 [Ex. 1095 at 1588-89]. Adair appealed, arguing the Board erred by (1) failing to assess material differences in view of the patent claim being copied, (2) establishing an absolute requirement that pre-critical date claims be patentable, (3) not applying applicable law, and (4) abusing its discretion in failing to consider other claims as pre-critical support for claim 24. The Federal Circuit affirmed the Board. *Adair v. Carter*, 101 U.S.P.Q.2d 1625, 1630 (Fed. Cir. 2012). Ex. 1095.

C. Level of Ordinary Skill in the Art

The invention's field involves humanizing non-human antibodies, e.g., mouse

monoclonal antibodies. A POSITA¹ would have held a Ph.D. or equivalent in chemistry, biological chemistry, structural biology or a closely related field, or an M.D. with practical academic or industrial experience in antibody development, including humanization of antibodies for therapeutic development and use in humans. See, e.g., Ex. 1003 at ¶¶26-28; Ex. 1004 at ¶¶38-39. Such a person would have the educational background above with experience related to antibody structural characterization and engineering. Id. Such experience can include three dimensional computer modeling of immunoglobulin structures, antibody domain and sequence manipulation and swapping, CDR grafting and framework substitution in humanizing antibodies, construction and expression of recombinant antibodies, antibody binding (specificity and affinity) testing, immunogenicity testing and the like. *Id.* Such person may have consulted with one or more team members of experienced professionals to develop a humanized monoclonal antibody for therapeutic use, including consulting with others to select non-human monoclonal antibodies (such as a mouse monoclonal antibody) for humanization, as well as subsequent testing of the humanized antibody and its intermediates. *Id.* Such a person would also have been well-versed in the

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¹ All references herein to the knowledge or understanding of a POSITA or a POSITA's interpretation or understanding of a prior art reference are as of the earliest possible priority date unless specifically stated otherwise.

world-wide literature that was available as of the priority date. *Id*.

D. Claim Construction

The '213 patent claims presumably possess their "broadest reasonable construction in light of the specification of the patent in which it appears." 37 C.F.R. § 42.100(b); *In re Cuozzo Speed Techs., LLC*, 793 F.3d 1268 (Fed. Cir. 2015) (affirming broadest reasonable construction standard in IPR). Under the broadest reasonable construction, a POSITA would understand the claim terms below at least include the following meanings.²

"A Humanized Antibody Variable Domain" (Claims 1, 62 and 80), "An antibody" (Claim 30) or "A Humanized Antibody" (Claim 63), "A Humanized Variant of a Non-Human Parent Antibody" (Claims 64 and 79) or "A Humanized Antibody Heavy Chain Variable Domain" (Claim 64). The independent claims of the '213 patent each contain a variation of the preamble phrase, "A Humanized Antibody" set forth above. A POSITA would understand "a humanized antibody" to include an antibody or antibody fragment that has been humanized, i.e., made more human-like. A POSITA would also understand that none of the claims relate to a

² Mylan does not concede that the claims can be construed to achieve reasonable certainty. Mylan explicitly does not waive any argument or invalidity position under 35 U.S.C. § 112, or any other invalidity position not presented herein.

single, specific antibody or antibody fragment. Even in claim 30, where the phrase "A humanized antibody" is modified with "which binds p185^{HER2}," the claim is not limited to a particular antibody.

"And Further Comprising a Framework Region (FR) Amino Acid Substitution at a Site Selected From the Group Consisting Of". Independent claims 1, 30, 62, 63, 66, 79 and 80 of the '213 patent include a Markush Group list of amino acid residues from which a framework region substitution is chosen. Markush Group members are accorded functional equivalency status for purposes of claim construction. See Ecolochem, Inc. v. Southern California Edison Co., 91 F.3d 169 (Fed.Cir. 1996) ("By claiming a Markush group ... members of the group are functionally equivalent" citing *Application of Skoll*, 523 F.2d 1392 (C.C.P.A. 1975)). As none of the claims are limited to a specific antibody, and all Markush Group members are functional equivalents of each other for the purpose of creating a humanized antibody, the broadest reasonable interpretation to a POSITA would be that any of the recited residues can be equally substituted for any given antibody. Thus, it is assumed for purposes of claim construction in this proceeding that each of the recited substitutions is available for humanization of an antibody.

"Numbering System Set Forth in Kabat". Independent claims 1, 30, 62, 63, 66, 79 and 80 of the '213 patent include the limitation "utilizing the numbering system set forth in Kabat." The '213 patent specifically ties its numbering system to

two references: "Kabat, E.A. et al., Sequences of Proteins of Immunological Interest (National Institutes of Health, Bethesda, Md.) (1987) and (1991)". See Ex. 1001 at col. 10, lns. 45-49. As noted above, the Kabat 1987 [Ex. 1052] and Kabat 1991 [Ex. 1055] data derives from a database of publicly available antibody sequences, formatted to display the sequences in alignment with each other and in a numerical sequence order. Kabat 1987 and 1991 also show boundaries of known antibody regions, including the three Complementarity Determining Regions (CDRs) and four Framework Regions (FRs) in each antibody chain variable domain. The broadest reasonable construction, "utilizing the numbering system set forth in Kabat," encompasses the Kabat 1987 and Kabat 1991 designations, including the amino acid residue positions set forth in Kabat, but also including the boundary designations for CDR and FR structures.

"Up To 3-Fold More". The '213 patent's claim 65 limits independent claim 79 further to a "humanized variant ... bind[ing] the antigen up to 3-fold more in the binding affinity than the parent antibody binds antigen." The broadest reasonable

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³ Dr. Padlan notes there are no significant differences between the Kabat 1987 and Kabat 1991 numbering systems, including CDR and FR boundary designations. Ex. 1003 at fn. 6. However, the priority document (U.S. Patent Application No. 07/715,272) only relies on Kabat 1987, and not Kabat 1991. *Id*.

interpretation of this claim includes all binding affinity values "up to" 3-fold more, *i.e.*, any value no matter how small and greater than zero "up to" 3-fold more.

E. Patents and Printed Publications Relied On In This Petition Petitioner relies on the following patents and printed publications:

1. EP 0403156 ("Kurrle") [Ex. 1071]

Kurrle, published on December 19, 1990, detailed the humanization of a mouse monoclonal antibody (BMA 031) against the human alpha/beta T-cell receptor. Ex. 1071 at Abstract. Kurrle provided guidance to a POSITA regarding further refinement of the variable domain FR region, making "essentially a human antibody with a much lower immunogenicity in patients." *See id.* at 3, ll. 8-12 ("A further refinement involves humanization of the variable regions. Only the complementarity determining regions and *selected framework amino acids necessary for antigen binding* are maintained murine. The remaining framework regions are converted to human sequences.") (Emphasis added).

Kurrle taught the four amino acids on either side of a CDR contribute to antibody binding:

"Molecular models of antibodies have shown that the actual CDR loops can contain amino acids up to 4 amino acids away from the 'Kabat' CDRs. Therefore, maintaining at least the major amino acid differences (in size or charge) within 4 amino acids of the CDRs as murine may be beneficial."

Kurrle [Ex. 1071] at 8, ll.27-29. Kurrle taught such "differences within 4 amino acids" should be "maintained murine." *Id.* at 8, ll.28-31. Kurrle further recommended using a simplified computer model based on sequence homology with solved antibody structures to judge the proximity of framework amino acid residues with the CDRs. *Id.* at 8, ll.32-36. Existing human framework residues could be switched to a consensus human residue at such positions. *Id.* at ll.38-46.

Applying one or more such criteria, Kurrle made four humanized versions of their antibody (CIV-1, CIV-2, CIV-3 and CIV-4), each time substituting select FR residues in the human antibody for the corresponding residue in the non-human (mouse) antibody. *See id.* at Tables 6A and 6B.⁴ Using their roadmap, Kurrle made several FR substitutions in the light and heavy chain, including at positions **4L**, **69H**, **71H**, **73H** and **76H**. *See* Ex. 1003 at Padlan Exhibit B, ¶142-58. The '213 patent claims these very residue substitutions. *Id*.

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⁴ Dr. Padlan notes that Kurrle did not use the Kabat numbering convention in Tables 6A and 6B for the antibody heavy chain. Ex. 1003 at fn 4. To follow '213 patent's numerical convention of the "numbering system set forth in Kabat," Dr. Padlan aligned the amino acid sequences in Table 6A (heavy chain) with the Kabat 1987 numbering system [Ex. 1052], as seen in Ex. 1003, Padlan Exhibit B.

2. Queen 1990 [Ex. 1050]

Queen 1990 is a PCT application filed December 28, 1989, and published July 26, 1990. Queen 1990 advanced Queen 1989's [Ex. 1034] methodology providing four explicit criteria for humanizing non-human antibodies. Criterion I of Queen 1990 relates to the choice of the acceptor human framework:

Criterion I: As acceptor, use a framework from a particular human immunoglobulin that is unusually homologous to the donor immunoglobulin to be humanized, or use a consensus framework from many human antibodies....

Ex. 1050 at 14, ll. 17-32.

Also like Queen 1989, Queen 1990 teaches that if a human FR residue is rare or unusual in humans, while the mouse residue is common (or conserved) in humans, substitute for the conserved mouse residue at that sequence position:

Criterion II: If an amino acid in the framework of the human acceptor immunoglobulin is unusual (i.e. "rare", which as used herein indicates an amino acid occurring at that position in no more than about 10% of human heavy (respectively light) chain V region sequences in a representative data bank), and if the donor amino acid at that position is typical for human sequences (i.e. "common", which as used herein indicates an amino acid occurring in at least about 25% of sequences in a representative data bank), then the donor amino acid rather than the acceptor may be selected....

Id. at 15, ll. 21-37. The prior art thus knew maintaining highly conserved residues was

important to minimize immunogenicity. Ex. 1003 at ¶116, 121.

Queen 1990 Criterion III also suggests substituting at CDR-adjacent positions:

Criterion III: In the positions immediately adjacent to one or more of the 3 CDR's in the primary sequence of the humanized immunoglobulin chain, the [mouse] donor amino acid(s) rather than acceptor amino acid may be selected. These amino acids are particularly likely to interact with the amino acids in the CDR's and, if chosen from the [human] acceptor, to distort the donor CDR's and reduce affinity. Moreover, the adjacent amino acids may interact directly with the antigen (Amit et al., Science, 233, 747-753 (1986), which is incorporated herein by reference) and selecting these amino acids from the [mouse] donor may be desirable to keep all the antigen contacts that provide affinity in the original antibody.

Id. at 16, Il. 1-12. Kabat and Chothia identified the CDR boundaries, both in sequence and structurally. Residues "immediately adjacent" to Kabat's CDRs are limited: 30H, 36H, 49H, 66H, 94H, 103H in the heavy chain; and 23L, 35L, 49L, 57L, 88L, and 98L in the light chain; residues "immediately adjacent" to Chothia's hypervariable regions include: 25L, 33L, 49L, 53L, 90L, 97L, 25H, 33H, 52H, 56H, 95H and 102H. '213 patent claims include 36H and 98L. Kabat 1987 [Ex. 1052]; Ex. 1003 at ¶¶159-69.

Queen 1990 placed further limitations on the molecular modeling criteria

Queen 1989 established, calling for pinpointing framework residues that possess an
atom that is within about 3Å of a CDR atom and thus likely to make a CDR contact:

Criterion IV: A 3-dimensional model, typically of the original [mouse] donor antibody, shows that certain amino acids outside of the CDR's are close to the CDR's and have a good probability of interacting with amino acids in the CDR's by hydrogen bonding, Van der Waals forces, hydrophobic interactions, etc. At those amino acid positions, the donor amino acid rather than the [human] acceptor immunoglobulin amino acid may be selected. Amino acids according to this criterion will generally have a side chain atom within about 3 angstrom units of some site in the CDR's and must contain atoms that could interact with the CDR atoms according to established chemical forces, such as those listed above. Computer programs to create models of proteins such as antibodies are generally available and well known to those skilled in the art.

Id. at 16, Il. 14-31 (citations omitted). Queen 1990 further teaches deriving these "contact" residues from known antibody structures. *Id.* Such framework residues are more likely to be important in influencing how CDRs interact with the antigen.

3. Furey [Ex. 1125]

Well prior to Queen and Kurrle's humanization efforts, Furey et al. established the structural importance of framework residues that established tight hydrogen bonding with hypervariable (CDR) residues, including at claimed position 66L in the light chain variable domain, to maintain CDR2 conformation. Ex. 1125 at Table 4. Furey therefore taught well before the alleged priority date that claimed V_L chain residue 66L contacted CDR2 residues via hydrogen bonds, and thus was a potential candidate for substitution according to the teachings of Kurrle and Queen 1990.

4. Chothia & Lesk [Ex. 1062]

Chothia & Lesk also established certain residues important for maintaining antibody structure, disclosing that "[t]he major determinants of the tertiary structure of the framework are the residues buried within and between the [V_L and V_H] domains," [Ex. 1062 at 4] summarized in Table 4 (reproduced below):

	V _L domains			V _H domains		
Position	Residues in known structures	A.S.A.* (Å ²)	Position	Residues in known structures	A.S.A.* (Ų)	
4	L,M	6	4	L	14	
6	ď.	12	6	$Q_{\cdot}E$	16	
19	Λ,	11	18	L	21	
21	I,M	ł	20	L	0	
23	C	0	22	(,	0	
25	G.A.S	13	24	S,V,T,A	8	
33	V.L	3	34	M.Y	4	
35	W	0	36	W.	0	
37	Q	30	38	R	13	
47	L,I.W	8	48	I,V	1	
48	I	24	49	A.G	0	
62	F	11	51	LV.8	4	
64	G.A	13	69	I.V,M	13	
71	A.F.Y	2	78	L,F	0	
73	$_{L,F}$	0	80	L	0	
75	I.V	0	82	M,L	0	
82	D	4	86	D	2	
84	A.S	11	88	A.G	3	
86	Y	0	90	Y	0	
88	C	0	92	Ċ.	0	
90	A.S.Q.N	7	104	G	11	
97	V,T,G	18	106	G	19	
99	\mathbf{G}	3	107	T.S	17	
101	G	11	109	Ÿ	2	
102	T	1			-	
104	L.V	2				

^a Mean accessible surface area (A.S.A.) of the residues in the Fab structures NEWM, MCPC603, KOL and J539 and in the V₁ structures REI and RHE.

Id. at 7, Table 4. These residues, which maintain tertiary structure (immunoglobulin chain interactions) of the framework, overlap with important CDR contact residues already disclosed in the prior art as well as known highly conserved residues, see Ex. 1003 at ¶¶137-39, fn 11, narrowing the list of substitutable residues significantly. Such residues—including claimed residues 4L, 62L, 73L, 4H, 36H, 69H, 78H and 92H—constitute potential substitution candidates under Kurrle and Queen 1990. Id.

5. Chothia 1985 [Ex. 1063]

Chothia 1985 disclosed "buried" residues involved in the "packing of the VL and VH β-sheets in the conserved 'framework'. . .." Chothia 1985 at Abstract [Ex. 1063 at 2]. "When the VL and VH domains pack together, residues from these edge strands form the central part of the interface and give what we call a three-layer packing; i.e. there is a third layer composed of side-chains inserted between the two backbone side-chain layers that are usually in contact. *The 12 residues that form the central part of the three observed VL-VH packings are absolutely or very strongly conserved in all immunoglobulin sequences*." *Id.* (emphasis added). One of the buried residues in the VL-VH interface disclosed by Chothia 1985 includes claimed residue **93H**. *See* Chothia 1985 at Table 4 [Ex. 1063].

6. Hudziak [Ex. 1021]

Hudziak published in March 1989, confirming p185^{HER2}'s role in carcinoma development. Ex. 1021 at Abstract. Hudziak had earlier-correlated p185^{HER2} gene amplification and carcinoma development, showing high p185^{HER2} levels correlated to negative prognoses and high relapse probability; and amplifying p185^{HER2} in vitro created resistance to cytotoxic (TNF-α) treatment. *Id.* Hudziak "prepared monoclonal antibodies against the extracellular domain of p185^{HER2}..." and chose "[o]ne monoclonal antibody (4D5)," which "was characterized in more detail and was shown to inhibit in vitro proliferation of human breast tumor cells overexpressing p185^{HER2} and, furthermore, to increase the sensitivity of these cells to the cytotoxic effects of

TNF- α ." *Id.* In growth inhibition studies, "[m]aximum inhibition was obtained with monoclonal antibody 4D5, which inhibited cellular proliferation by 56%." *Id.* (emphasis added). Hudziak confirmed "the combination of TNF- α and monoclonal antibody 4D5 reduced the [listed] tumor cell number to a level below that initially plated," and "indicated the induction of a cytotoxic response." Ex. 1021 at 6.

Monoclonal antibody		Relative cell proliferation*		
7C2	79.3	±	2.2	
2C4	79.5	±	4.4	
7D3	83.8	±	5.9	
4D5	44.2	±	4.4	
3E8	66.2	±	2.4	
6E9	98.9	±	3.6	
7F3	62.1	±	1.4	
3H4	66.5	±	3.9	
2H11	92.9	±	4.8	
40.1.H1	. 105.8	±	3.8	
4F4	94.7	±	2.8	

Hudziak, Table 1 [Ex. 1021 at 4]. Hudziak concluded that "[m]onoclonal antibodies specific for p185^{HER2} may therefore be useful therapeutic agents for the treatment of human neoplasias, including certain mammary carcinomas, which are characterized by the overexpressing of p185^{HER2}." Ex. 1021 at 7.

F. The Prior Art Renders The Challenged Claims Obvious

1. Detailed Instructions for Humanizing Antibodies Were Widely Available Before the '213 Patent Filing

Multiple research institutions—including Genzyme Corp. [Ex. 1071], Protein Design Labs [Ex. 1050], the Medical Research Council and the National Institutes of Health, amongst others—published before the '213 patent's filing date efforts to

humanize antibodies to avoid the immunogenic reactions observed with non-human monoclonal antibody therapeutics. *See* Ex. 1071 at 3, Il. 8-12; Ex. 1050 at Abstract; Ex. 1003 at ¶94; Ex. 1004 at ¶50. The field recognized that earlier efforts (*e.g.*, chimeric antibodies, CDR grafting) often resulted in non- or poor binding, with immunogenicity remaining a concern. *See* Ex. 1050 at 5, Il. 30-33; Ex. 1073 at 9, Il. 12-19; Ex. 1003 at ¶89-92; Ex. 1004 at ¶46-47.

Queen 1990 detailed the importance of preserving certain mouse framework positions in the resulting humanized antibody in order to maintain CDR conformation and antigen binding. Ex. 1050 at 16, ll. 2, 14–15. The prior art, thus, already provided detailed pathways to humanize antibodies for therapeutic use which would "be substantially non-immunogenic and retain substantially the same affinity as the donor immunoglobulin to the antigen." *See id.* at Abstract; Ex. 1003 at ¶¶92-107.

Kurrle used logic similar to Queen's, replacing several human FR sites with mouse residues within the variable region of the light and heavy chains. Ex. 1071 at Tables 6A and 6B; Ex. 1003 at ¶108-111.

Simply put, many scientific research groups were making "humanized antibodies" more than a year prior to the '213 patent's earliest filing date and publishing detailed instructions for doing the same. The copious prior art demonstrates that modification and humanization as claimed in each challenged claim was not only anticipated, but plainly obvious.

G. <u>Ground 1</u>: Claims 1, 2, 25, 29, 63, 66, 71, 75, 76, 78, 80 and 81 Are Unpatentable As Anticipated By Kurrle

1. Independent Claim 1 is Anticipated by Kurrle

Independent claim 1 of the '213 patent recites "[a] humanized antibody variable domain comprising," the elements (1) "non-human Complementarity

Determining Region (CDR) amino acid residues which bind an antigen incorporated into a human antibody variable domain," and (2) FR substitutions at "a site selected from the group consisting of: 4L, 38L, 43L, 44L, 58L, 62L, 65L, 66L, 67L, 68L, 69L, 73L, 85L, 98L, 2H, 4H, 36H, 39H, 43H, 45H, 69H, 70H, 74H, and 92H, utilizing the numbering system set forth in Kabat."

The "humanized antibody variable domain" element is disclosed in Kurrle, which taught "humanised and civilised versions of [mouse monoclonal] antibodies." Ex. 1071 at Abstract; Ex. 1003 at ¶¶108-111, 143. Kurrle also disclosed "non-human Complementarity Determining Region (CDR) amino acid residues which bind an antigen" and "a Framework Region (FR) amino acid substitution" incorporated into a human antibody variable domain, referring to the "civilised" antibodies as those where "[o]nly the *complementarity determining regions* and *selected framework amino acids* necessary for antigen binding are maintained *murine*." Ex. 1071 at 3, Il. 9-12 (emphasis added); Ex. 1003 at ¶143.

Kurrle substituted several corresponding murine amino acids for human framework residues under Kabat's numbering system, including **4L** and **69H**, as

found in claim 1. *See* Ex. 1071 at 25, 26, Tables 6A and 6B; Ex. 1003 at ¶¶142-145, Padlan Exhibit B. Claim 1 is anticipated.

2. Kurrle Anticipates Dependent Claims 2, 25 and 29

Claim 2: Claim 2 depends on claim 1, and further recites, "wherein the substituted residue is the residue found at the corresponding location of the non-human antibody from which the non-human CDR amino acid residues are obtained." This is precisely what Kurrle did. See Ex. 1071 at 8, Il. 45-47 ("In one position (#93) the human consensus sequence is the same as [in the mouse sequence]. One could rationalize changing [the human acceptor antibody residue] back to [mouse], so this change was incorporated..."). This is a basic step in the humanization process as taught by Kurrle. See Ex. 1003 at ¶146. Claim 2 is thus also anticipated by Kurrle.

<u>Claim 25</u>: Claim 25 depends on claim 1, and further recites "wherein the residue at site 69H has been substituted." Because framework residue **69H** was substituted with the murine residue in Kurrle's humanized anti-T-cell receptor antibody, *see* claim 1 (§VI.G.1), Kurrle anticipates claim 25. Ex. 1003 at ¶147.

<u>Claim 29</u>: Claim 29 also depends on claim 1, and further recites "[a]n antibody comprising the humanized variable domain of claim 1." Kurrle's explicit goal was to create an antibody comprising the humanized variable domain: "The resulting mAb of the present invention is thus essentially a human antibody with a much lower

immunogenicity in patients." Ex. 1071 at 3, ll. 9-12; *see also* 2, ll. 2-4; Ex. 1003 at ¶148. Kurrle anticipates Claim 29.

3. Independent Claim 63 is Anticipated by Kurrle

Claim 63 of the '213 patent is drawn to an antibody with structural components substantially identical to those of claim 29, *i.e.*, the same "humanized antibody" incorporating the same claimed non-human CDRs and completely overlapping substituted framework residues as in claim 1. *See* §VI.G.2, *supra*. Accordingly, because the structural components are the same, the same <u>function</u> (*i.e.*, "which lacks immunogenicity compared to a non-human parent antibody upon repeated administration to a human patient in order to treat a chronic disease in that patient") is also present. *See Atlas Powder Co. v. Ireco Inc.*, 190 F.3d 1342, 1347 (Fed. Cir. 1999) ("'[T]he discovery of a previously unappreciated property of a prior art composition, or of a scientific explanation for the prior art's functioning, does not render the old composition patentably new to the discoverer."); Ex. 1003 at ¶¶149-51.

Not only is this an inherent aspect of the claimed humanized antibodies, this is in fact an explicitly stated goal of all antibody humanization projects. *See* Ex. 1071 at 3, ll. 8-12 ("A further refinement involves humanization of the variable regions ...

[T]he resulting mAb of the present invention is thus essentially a human antibody with a much lower immunogenicity in patients."); Ex. 1003 at ¶¶150-51. One of ordinary skill in the art would thus know that Kurrle's humanized antibodies would

also "lack immunogenicity compared to a non-human parent antibody upon repeated administration". Claim 63 is anticipated.

4. Independent Claim 66 and Dependent Claims 71, 72, 75 and 76 are Anticipated by Kurrle

Independent claim 66 shares elements with claims 1 and 63, which are met as demonstrated above. *See* §§ VI.G.1 & 3, *supra*; Ex. 1003 at ¶152-53. Claim 66's substitutable amino acid residues are "selected from the group consisting of: 24H, 73H, 76H, 78H, and 93H," under Kabat's numbering system. As Kurrle substituted residues **73H** and **76H**, Ex. 1003 at Padlan Exhibit B and ¶152-53, it anticipates claim 66.

Dependent claims 71, 72, 75 and 76 recite "wherein the residue at site 73H has been substituted" (claim 71), "wherein the residue at site 76H has been substituted" (claim 72), "which further comprises an amino acid substitution at site 71H" (claim 75), and "which further comprises amino acid substitutions at sites 71H and 73H" (claim 76). Kurrle disclosed the substitution of amino acid residues **71H**, **73H** and **76H** in their humanized anti-T-cell receptor monoclonal antibody. *See* Ex. 1071 at Table 6B; Ex. 1003 at Padlan Exhibit B and ¶54. Accordingly, and in view of the discussion for claims 1 and 66, *see* §§VI.G.1 & 3, *supra*; Ex. 1003 at ¶¶142-45, 152-54, Kurrle anticipates dependent claims 71, 72, 75 and 76.

5. Independent Claim 80 and Dependent Claim 81 Are Anticipated by Kurrle

Claim 80: Independent claim 80 recites "[a] humanized antibody variable domain comprising non-human Complementarity Determining Region (CDR) amino acid residues which bind an antigen incorporated into a human antibody variable domain, and further comprising a Framework Region (FR) amino acid substitution." Claim 80 further recites the "substituted FR residue: (a) noncovalently binds antigen directly; (b) interacts with a CDR; or (c) participates in the V_L - V_H interface by affecting the proximity or orientation of the V_L and V_H regions with respect to one another...." Claim 80 then recites a list of substitutable amino acid residues that differ from claim 1 by adding amino acid residues 73H, 76H, 78H and 93H to the list. As with claims 1 and 63, residues 4L and 69H, and additional residues 73H and 76H are substituted in Kurrle. §§VI.G.1 & 3, supra.

The additional recited elements, which are noted functions of the substituted residues, do not add anything new to the claim. *See* claim 63, §VI.G.3; Ex. 1003 at ¶¶155-56; *see also Atlas Powder*, 190 F.3d at 1347. Even if the inherency of these functions were discounted (they should not be), Kurrle explicitly teaches interaction of the framework residues with the CDR as a reason for substitutability. *See* Ex. 1071 at 8, Il. 28-29 and 32-40 (use of a "simplified computer model" to determine whether or not FR residues were close enough to CDRs to influence binding); Ex. 1003 at ¶¶108-111, 156. Accordingly, Kurrle at least teaches substitution of a framework

residue that "interacts with a CDR," *i.e.*, limitation "(b)" from claim 80, and therefore anticipates claim 80.

<u>Claim 81</u>: Claim 81 depends on claim 80, and further recites, "wherein the substituted residue is the residue found at the corresponding location of the non-human antibody from which the non-human CDR amino acid residues are obtained." This is taught by Kurrle. *See* §VI.G.2, *supra*; Ex. 1071 at Tables 6A and 6B; Ex. 1003 at ¶155-58. Claim 81 is thus also anticipated by Kurrle.

H. <u>Ground 2</u>: Claims 1, 2, 4, 29, 62, 63, 64, 65, 80 and 81 are Anticipated by Queen 1990

1. Independent Claim 1 is Anticipated by Queen 1990

The first part of claim 1, "[a] humanized antibody variable domain," is disclosed in Queen 1990. Queen's stated goal was creating "a humanized antibody variable domain" by not only swapping CDRs, but also manipulating the framework region of the variable domain, as claim 1 of the '213 patent recites. Queen explicitly provides for "novel methods for designing humanized immunoglobulins having one or more complementarity determining regions (CDR's) from a donor immunoglobulin and a framework region from a human immunoglobulin...." Ex. 1050 at Abstract; Ex. 1003 at ¶160. Queen 1990 provides a detailed roadmap with specific criteria used in designing humanized immunoglobulins. *Id.* at 14, Il.10-11; Ex. 1003 at ¶160-69, 143-51. Queen 1990 emphasized the importance of framework positions adjacent to the CDR: "Each humanized immunoglobulin chain may comprise about 3 or more

amino acids from the donor immunoglobulin in addition to the CDR's, *usually at least one of which is immediately adjacent to a CDR in the donor immunoglobulin.*" Ex. 1050 at Abstract. The POSITA can readily envision such locations. *See* Ex. 1003 at ¶¶164-66.

Queen 1990 encapsulated this rule in Criterion III, which states:

In the positions immediately adjacent to one or more of the 3 CDR's in the primary sequence of the humanized immunoglobulin chain, the donor [mouse] amino acid(s) rather than acceptor [human] amino acid may be selected. These amino acids are particularly likely to interact with the amino acids in the CDR's and ...[m]oreover, the adjacent amino acids may interact directly with the antigen ... and selecting these amino acids from the donor may be desirable to keep all the antigen contacts that provide affinity in the original antibody.

Ex. 1050 at 16, ll. 1-12 (emphasis added, citations omitted); Ex. 1003 at ¶164.

Mylan's expert, Dr. Padlan, explained that "one of ordinary skill in the art at the time of the '213 patent . . . would have readily understood that Queen 1990 (specifically Criterion III) explicitly taught the substitution of framework sites **immediately adjacent** to CDRs." Ex. 1003 at ¶165. Using the numbering system set forth by Kabat 1987,⁵ the "immediately adjacent" framework residues to CDRs as

⁵ While Dr. Padlan uses the Kabat 1987 reference for designating the amino acid positions according to the Kabat numbering system, there were "no meaningful

taught by Queen 1990 and recited in claim 1 include <u>98L</u> and <u>36H</u>. See Ex. 1003 at Padlan Exhibit C and ¶¶159-69; Section VI.E.2, supra.

Thus, Queen 1990's explicit teaching to substitute CDR-adjacent framework region amino acid positions would inevitably include substitutions at the claimed amino acid residues of **98L** and **36H**. Queen 1990 thus anticipates claim 1.

2. Queen 1990 Anticipates Dependent Claims 2, 4 and 29

Claim 2: Claim 2's additional limitation "wherein the substituted residue is the residue found at the corresponding location of the non-human antibody from which the non-human CDR amino acid residues are obtained" is also disclosed by Queen 1990. See Ex. 1050 at 7, 1. 36-8, 1. 1 ("substitutions of a human framework amino acid of the acceptor (*i.e.*, human) immunoglobulin with a corresponding amino acid from a donor (*i.e.*, non-human) immunoglobulin"); Ex. 1003 at ¶170. Queen 1990 anticipates claims 2, 4, and 29.

<u>Claim 4</u>: Claim 4 depends on claim 1, and further recites "wherein the human antibody variable domain is a consensus human variable domain." Queen 1990 expressly teaches this by disclosing in Criterion I that "[a]s acceptor, ... use *a*

differences in the Kabat numbering system, including the CDR boundaries, between Kabat 1987 and Kabat 1991." Ex. 1003 at fn. 5.

consensus framework from many human antibodies." See Ex. 1050 at 14, ll.17-20 (Criterion I); Ex. 1003 at ¶¶119, 171. Queen 1990 anticipates claim 4.

Claim 29: Claim 29 depends on claim 1, and further recites "[a]n antibody comprising the humanized variable domain of claim 1." As Dr. Padlan explains, the goal of antibody humanization programs was to create a humanized variable domain. See, e.g., Ex. 1050 at 6, Il. 21-25 ("mouse complementarity determining regions, with or without additional naturally-associated mouse amino acid residues, can be used to produce human-like antibodies"); Ex. 1003 at ¶172. A POSITA would thus recognize that Queen's aim was to create therapeutic-quality antibodies with a humanized variable domain in order to maintain a high level of binding and affinity. Ex. 1003 at ¶172. Queen 1990 anticipates claim 29.

3. Independent Claim 62 is Anticipated by Queen 1990

Claim 62 shares claim 1's FR substitutable residues list, but adds residues 46L, 75H, 76H and 78H. As discussed above for claim 1, *see* §VI.H.1, *supra*, Queen 1990 discloses residues **98L** and **36H** as also inevitably requiring substitution.

Claim 62 also differs from claim 1 by adding the phrase, "incorporated into a consensus human variable domain." Ex. 1003 at ¶173. Queen 1990 also disclosed in Criterion I that "[a]s acceptor, ... use *a consensus framework* from many human antibodies." *See* Ex. 1050 at 14, ll. 17-20; Ex. 1003 at ¶174; §VI.H.2, *supra*. Queen 1990 thus anticipates claim 62 of the '213 patent.

4. Independent Claim 63 is Anticipated by Queen 1990

Claim 63 differs from claim 62 (incorporating into the claimed "antibody" the "humanized variable domain" comprising the recited CDRs and substituted FR residues from claim 1) by further describing the claimed humanized antibody as lacking "immunogenicity compared to a non-human parent antibody upon repeated administration to a human patient."

As above, this is a non-patentable distinction. *See* §VI.G.3, citing to *Atlas Powder*, 190 F.3d 1342; Ex. 1003 at ¶176. Queen 1990 explicitly taught this goal: "When combined into an intact antibody, the humanized immunoglobulins of the present invention *will be substantially non-immunogenic* in humans..." Ex. 1050 at Abstract; Ex. 1003 at ¶176-77. Claim 63 is also anticipated by Queen 1990.

5. Independent Claim 64 is Anticipated by Queen 1990

Claim 64 recites "a humanized variant of a non-human parent antibody which binds an antigen; comprising a human variable domain comprising the most frequently occurring amino acid residues at each location in all human immunoglobulins of a human heavy chain immunoglobulin subgroup; wherein amino acid residues forming Complementarity Determining Regions (CDRs) thereof comprise non-human antibody amino acid residues, and further comprises a Framework Region substitution where the substituted FR residue: (a) noncovalently binds antigen directly; (b) interacts with a CDR; (c) introduces a glycosylation site

which affects the antigen binding or affinity of the antibody; \underline{or} (d) participates in the V_L - V_H interface by affecting the proximity or orientation of the V_L and VH regions with respect to one another." (Emphasis added).

Queen 1990 anticipates claim 64. As with claims 1, 4 and 29, Queen 1990 disclosed an antibody incorporating a humanized variable domain with a consensus sequence (*i.e.*, "most frequently occurring amino acid residues at each location in all human immunoglobulins of a human heavy chain immunoglobulin subgroup"). *See* §§VI.H.2 & 3, *supra*; Ex. 1003 at ¶¶183-85; Ex. 1050 at 14, ll. 17-20 ("As acceptor, ... use *a consensus framework* from many human antibodies.").

While the remaining limitations are merely stated functions of the humanized antibody, *see* §VI.G.3, *supra*, Queen 1990 also disclosed at least functions (a) and (b) above in Criterion III:

[I]mmediately adjacent ... amino acids are *particularly likely to interact* with the amino acids in the CDR's and, if chosen from the acceptor, distort the donor CDR's and reduce affinity. Moreover, the adjacent amino acids may interact directly with the antigen ... and selecting these amino acids from the donor may be desirable to keep all the antigen contacts that provide affinity in the original antibody.

Ex. 1050 at 16, ll. 1-12 (emphasis added); Ex. 1003 at ¶185. Because Queen 1990 teaches one to substitute "immediately adjacent" residues **98L** and **36H**, *see* §VI.H.1 *supra*, and because Queen 1990 teaches that those residues "are particularly likely to

interact with the amino acids in the CDR's and ... may interact directly with the antigen," Queen 1990 anticipates claim 64. Ex. 1003 at ¶¶1183-85.

6. Claims 80 and 81 are Anticipated by Queen 1990

Claim 80: Claim 80 is also anticipated by Queen 1990. As discussed with claims 1 and 64, Criterion III of Queen 1990 explicitly teaches the selection of framework residues immediately adjacent to CDRs for substitution − this would include claimed residues 36H and 98L. See §§VI.H.1 & 5, supra; Ex. 1003 at Padlan Exhibit C and ¶¶ 186-89 (citing Ex. 1050 at 16, ll. 4-8). Queen 1990 explains that "selecting these amino acids from the donor may be desirable to keep all the antigen contacts that provide affinity in the original antibody." Ex. 1050 at 16, ll. 10-12; Ex. 1003 at ¶187.

Moreover, Criterion IV in Queen 1990 teaches "interact[ion] with a CDR" by disclosing that "certain amino acids outside of the CDR's are close to the CDR's and have a good probability of interacting with amino acids in the CDR's by hydrogen bonding, Van der Waals forces, hydrophobic interactions, etc." Ex. 1050 at 16, ll. 15-19; Ex. 1003 at ¶188. Given that, Queen 1990 anticipates Claim 80.

Claim 81: Claim 81 (which depends on claim 80) is also taught by Queen 1990, which disclosed "substitutions of a human framework amino acid of the acceptor (*i.e.*, human) immunoglobulin with a corresponding amino acid from a donor (*i.e.*, non-human) immunoglobulin." See Ex. 1050 at 7, 1. 36-8, 1. 1; Ex. 1003 at ¶190.

Claim 81 is also anticipated by Queen 1990.

7. Dependent Claim 65 is Anticipated by Queen 1990

Claim 65 depends from claim 79, further reciting that the humanized variant "binds the antigen up to 3-fold more in the binding affinity than the parent antibody binds antigen." Claim 65 is also taught by Queen 1990, which states "affinity levels can vary . . . and may be within about 4 fold of the donor immunoglobulin's original affinity to the antigen." See Ex. 1050 at 8, ll. 26-28 (emphasis added). Queen 1990 thus explicitly taught that a humanized antibody would have been expected to be "within about 4-fold," i.e. "up to 3-fold" in affinity as the original mouse antibody. Ex. 1033 at ¶¶178-82. Queen 1990 anticipates Claim 65.

- I. <u>Ground 3</u>: Claims 1, 2, 4, 25, 29, 62-67, 69, 80 and 81 Are Unpatentable As Obvious over Queen 1990 and Kurrle
 - 1. Claim 1 is Obvious Over Queen 1990 and Kurrle

Queen 1990 disclosed to a POSITA a detailed pathway for humanizing non-human monoclonal antibodies, with the expectation that the antibodies "will be substantially non-immunogenic in humans and retain substantially the same affinity as the donor immunoglobulin to the antigen ...," including:

• Criterion I: Choose an acceptor human framework antibody, including one that is "unusually homologous to the donor immunoglobulin to be humanized, or use a consensus framework from many human antibodies." Ex. 1050 at 14, 1. 17 – 15, 1. 20; Ex. 1003 at ¶¶119, 194;

- Criterion II: Once the human antibody is selected, evaluate whether amino acid residues in the framework of the human acceptor antibody are "rare" amongst human antibodies. If the residue is "rare" and the donor [mouse] antibody is more "typical for human sequences," choose the donor residue. Criterion II "helps ensure that an atypical amino acid in the human framework does not disrupt the antibody structure." Ex. 1050 at 15, Il. 22-37; Ex. 1003 at ¶¶120-21, 194;
- Criterion III: "In the positions immediately adjacent to the 3 CDR's in the humanized immunoglobulin chain, the donor [mouse] amino acid rather than acceptor [human] amino acid may be selected." Ex. 1050 at 15, ll. 1-12; Ex. 1003 at ¶122, 194; and
- Criterion IV: Generate a 3-dimensional model of the original donor antibody, and select amino acid positions where:

[C]ertain amino acids outside of the CDR's are close to the CDR's and have a good probability of interacting with amino acids in the CDR's Amino acids according to this criterion will generally have a side chain atom within about 3 angstrom units of some site in the CDR's and must contain atoms that could interact with the CDR atoms according to established chemical forces, such as those listed above.

Ex. 1050 at 16, l. 14 - 17, l. 2; Ex. 1003 at ¶¶123, 194.

Queen 1990 concludes that when the humanized variable regions are "combined into an intact antibody, the humanized light and heavy chains of the present invention will be substantially non-immunogenic in humans and retain substantially the same affinity as the donor immunoglobulin to the antigen...." Ex. 1050 at 8, Il. 21-26; Ex. 1003 at ¶195. Queen 1990 thus provided "the motivation along with a detailed roadmap for production of a humanized monoclonal antibody that can be used in human therapeutics." Ex. 1003 at ¶195.

Kurrle employed a similarly detailed roadmap to obtain a "humanized antibody variable domain" as claimed in claim 1, including the steps of: choosing the most similar human acceptor sequence (Criterion I of Queen 1990; *see* Ex. 1071 at 8; ll. 16-18), accounting for the adjacent residue rules of Queen 1990 (Criterion III of Queen 1990; *see* Ex. 1071 at 8, ll. 25-31), substituting CDR-contact residues using computer models based on solved structures (Criterion IV of Queen 1990; *see* Ex. 1071 at 8, ll. 32-36) and substituting "rare" amino acids in the human acceptor framework for "common" (consensus) amino acid residues (Criterion II of Queen 1990; *see* Ex. 1071 at 8, ll. 36-40). Ex. 1003 at ¶108-111, 196.

Using these guidelines, Kurrle made a total of 13 substitutions in the light chain framework region and 18 substitutions in the heavy chain framework region according to the Kabat numbering system, including claimed residues **4L** and **69H**.

See §§VI.E.1 & G.1, supra; Ex. 1003 at Padlan Exhibit B, ¶¶142-45, 197.

The combination of Queen 1990 and Kurrle thus provided ample motivation and a reasonable expectation of success that a humanized monoclonal antibody could be obtained with "a much lower immunogenicity in patients", Ex. 1071 at 3, ll. 11-12, while maintaining the binding affinity and specificity of the donor monoclonal antibody, and targeted the very species residues satisfying the claim 1 genus. Claim 1 is obvious over Queen 1990 in view of Kurrle. Ex. 1003 at ¶¶194-98.

2. Claims 2, 25 and 29 are Obvious Over Queen 1990 and Kurrle

Claim 2: Claim 2 is also taught by Queen 1990 and Kurrle. As discussed, this is a basic step in humanization, followed by many in the field, including Queen (Ex. 1050 at 7, 1. 36-8, 1. 1) and Kurrle (Ex. 1071 at 8, 1l. 28-29). See §VI.G.2 supra; Ex. 1003 at ¶199. Claim 2 is obvious over Queen 1990 and Kurrle.

Claim 25: Claim 25 depends on claim 1, and further recites "wherein the residue at site 69H has been substituted." Residue 69H was substituted in Kurrle's humanized anti-T-cell receptor antibody. See §§VI.G.1 & 2, supra; Ex. 1003 at ¶200. Accordingly, claim 25 is also obvious over Queen 1990 and Kurrle.

<u>Claim 29</u>: Claim 29 depends on claim 1, and further recites "[a]n antibody comprising the humanized variable domain of claim 1." The explicit goals of Queen 1990 and Kurrle was to create antibodies comprising a humanized variable domain: "the humanized light and heavy chains of the present invention will be substantially non-immunogenic in humans and retain substantially the same affinity as the donor

immunoglobulin to the antigen." Ex.1050 (Queen 1990) at 8, ll. 21-26; *see also* Ex. 1071 (Kurrle) at 3, ll. 26-28 and 2, ll. 2-4; Ex. 1003 at ¶201. Claim 29 is also obvious over Queen 1990 in view of Kurrle.

3. Claim 4 is Obvious Over Queen 1990 and Kurrle

Claim 4, which depends from claim 1, recites: "wherein the human antibody variable domain is a consensus human variable domain." Queen 1990 teaches the use of a human consensus variable domain as the human acceptor framework antibody, see Ex. 1050 at 14, ll. 17-20 ("As acceptor ... use a consensus framework from many human antibodies."), which would have motivated a POSITA to use the human "acceptor" framework together with the humanization methods of Kurrle. Ex. 1003 at \$\$\\$119, 202. Claim 4 is also obvious over Queen 1990 and Kurrle.

4. Claim 62 is Obvious Over Queen 1990 and Kurrle

Claim 62 differs from claim 1 by adding that the human variable domain is a "consensus human variable domain." *See* §§VI.B.1 & I.1, *supra*. Queen 1990 discloses the use of a consensus human variable domain in Criterion I of its humanization roadmap. Ex. 1050 at 14, Il. 17-20 ("As acceptor, ... use *a consensus framework* from many human antibodies."); Ex. 1003 at ¶¶206-207. Queen 1990 and Kurrle provided both the motivation and a reasonable expectation of success to make and use the remaining limitations, including substituting at claimed positions **98L** and **36H** (Ex. 1050; §H.1) and **4L**, **69H** and **76H** (Ex. 1071; §G.1). Ex. 1003 at ¶207.

Claim 62, as for claims 1 and 4 (see §§VI.I.1 & 2, supra) of the '213 patent is obvious over Queen 1990 and Kurrle.

5. Claim 63 is Obvious Over Queen 1990 and Kurrle

Claim 63 is similar to claim 62 (incorporating the CDRs and substituted FR residues of claim 1), and adds that the claimed humanized antibody "lacks immunogenicity compared to a non-human parent antibody upon repeated administration to a human patient to treat a chronic disease in that patient." This is the stated goal of all humanization projects, including that of Queen 1990 and Kurrle. *See* Ex. 1050 at Abstract ("the humanized immunoglobulins of the present invention *will be substantially non-immunogenic* in humans..."); Ex. 1071 at 3, Il. 11-12 ("The resulting mAb of the present invention is thus essentially a human antibody with a much lower immunogenicity in patients."); Ex. 1003 at ¶208-211. Claim 63 is also obvious over Queen 1990 and Kurrle.

6. Claim 64 is Obvious Over Queen 1990 and Kurrle

Queen 1990 and Kurrle also disclose the limitations of claim 64. Queen 1990 discloses an antibody incorporating a humanized variable domain comprising a consensus sequence. *See* §§VI.H.2 & 5, *supra*; Ex. 1050 at 14, ll. 17-20 ("As acceptor, ... use *a consensus framework* from many human antibodies."); Ex. 1003 at ¶¶212-215. Both Queen 1990 and Kurrle also taught humanized antibodies containing a non-human CDR and substituted FR residues. *See, e.g.*, Ex. 1071 (Kurrle) ("Only

the complementarity determining [sic] regions and selected framework amino acids necessary for antigen binding are maintained murine. The remaining framework regions are converted to human sequences."); Ex. 1003 at ¶213.

While the remaining limitations are merely stated functions of the humanized antibody, *see* §§VI.G.3, & H.5 *supra*, both Queen 1990 and Kurrle disclosed that certain framework residues were important because of their proximity to neighboring CDRs: "These amino acids are *particularly likely to interact with the amino acids in the CDR's* and, if chosen from the acceptor, distort the donor CDR's and reduce affinity." *See* Ex. 1050 at 16, ll. 1-12 (emphasis added); *see also* Ex. 1071 at 8, ll. 27-29; Ex. 1003 at ¶213. Queen 1990 and Kurrle provided the motivation and reasonable expectation of success to make the claimed "humanized variant of a non-human parent antibody." Claim 64 is obvious over Queen 1990 and Kurrle.

7. Claim 65 is Obvious Over Queen 1990 and Kurrle

Claim 65 depends from claim 79 and further recites that the humanized variant "binds the antigen up to 3-fold more in the binding affinity than the parent antibody binds antigen." The broadest reasonable interpretation of this limitation includes <u>any</u> increase in binding affinity "up to" the upper limit of "3-fold more," *i.e.*, <u>any</u> amount greater than 1-fold (*i.e.*, the parent binding affinity) and "up to 3-fold more." §VI.D, *supra*; Ex. 1003 at ¶217-220.

Dr. Padlan explained that to a POSITA, "it was the expectation when

humanizing antibodies . . . that a similar affinity, *i.e.*, slightly better or worse, would be obtained as compared to the parent (mouse) antibody. Thus . . . it would not have been unexpected that at least a moderate improvement in affinity would be achieved when humanizing some antibodies." Ex. 1003 at ¶218. Dr. Padlan further explains that "it was not unexpected that in this process, one could go beyond the parent antibody's original affinity, *i.e.*, an increase in affinity as claimed in claim 65." Ex. 1003 at ¶219. This is within the stated purpose of humanization, and thus any increase, including small and moderate increases incorporated within the scope of the claim, would have been expected, as stated by Queen 1990. Ex. 1050 at 8, Il.26-28 ("[A]ffinity levels can vary . . . and may be within about 4 fold of the donor immunoglobulin's original affinity to the antigen."); Ex. 1003 at ¶1216-220. For these reasons, claim 65 is also obvious over the combination of Queen 1990 and Kurrle.

8. Claim 66 is Obvious Over Queen 1990 and Kurrle

Both Queen 1990 and Kurrle disclose the claimed "humanized antibody heavy chain variable domain comprising non-human Complementarity Determining Region (CDR) amino acid residues which bind antigen incorporated into a human antibody variable domain," which is also essentially recited in claims 1 and 62. *See* §§VI.I.1 & 4, *supra*. Claim 66 further requires the framework substitution of one or more residues, *e.g.*, 24H, 73H, 76H, 78H and 93H. Kurrle, using Queen 1990's roadmap, substituted FR amino acids at claimed positions **73H** and **76H**, rendering the

humanized antibody "essentially a human antibody with a much lower immunogenicity in patients." Ex. 1071 at 3, ll. 11-12; Ex. 1003 at ¶221-22.

Both Queen 1990 and Kurrle provide the motivation and a reasonable expectation of success to make "a humanized antibody variable domain" as claimed in claim 66. Ex. 1003 at ¶222. Claim 66 is also obvious over Queen 1990 in view of Kurrle.

9. Claims 67, 71, 72, 75, 76 and 78 are Obvious Over Queen 1990, and Kurrle

Claim 67: Claim 67, which depends from claim 66, recites "wherein the substituted residue is the residue found at the corresponding location of the non-human antibody from which the non-human CDR amino acid residues are obtained." Both Queen 1990 and Kurrle disclosed this additional limitation. See, e.g., Ex. 1050 at 7, l. 36-8, l. 1 (disclosing "substitutions of a human framework amino acid of the acceptor (i.e., human) immunoglobulin with a corresponding amino acid from a donor (i.e., non-human) immunoglobulin."); Ex. 1003 at ¶223. Claim 67 is also obvious over Queen 1990 and Kurrle.

<u>Claims 71, 72, 75 and 76</u>: Claims 71, 72, 75 and 76, which all depend from claim 66, recite "wherein the residue at site 73H has been substituted" (claim 71), "wherein the residue at site 76H has been substituted" (claim 72) "which further comprises an amino acid substitution at site 71H" (claim 75), and "which further comprises amino acid substitutions at sites 71H and 73H" (claim 76). Kurrle

substituted the murine amino acid residues at claimed positions **71H**, **73H** and **76H**. Ex. 1003 at ¶224. Together with claim 66, §VI.I.8 *supra*, dependent claims 71, 72, 75 and 76 are also obvious over Queen 1990 in view of Kurrle.

Claim 78: Claim 78, which depends on claim 66, recites an antibody "comprising the humanized variable domain of claim 66." The goal of humanization methods, including Queen 1990 and Kurrle, was to create a therapeutic antibody comprising a humanized variable domain: "When combined into an intact antibody, the humanized light and heavy chains of the present invention will be substantially non-immunogenic in humans and retain substantially the same affinity as the donor immunoglobulin to the antigen." See Ex. 1050 at 8, ll. 21-26; Ex. 1071 at 3, ll. 26-28; Ex. 1003 at ¶227. Claim 78 is obvious over Queen 1990 and Kurrle.

10. Claim 69 is Obvious Over Queen 1990 and Kurrle

Claim 69 is dependent on claim 66, and further recites that, "the human antibody variable domain is a consensus human variable domain." Queen 1990 explicitly teaches using a consensus sequence as the human acceptor framework antibody: "As acceptor ... use a consensus framework from many human antibodies." Ex. 1050 at 14, ll. 17-20; Ex. 1003 at ¶228. In view of claim 66, see §VI.I.8, supra, claim 69 is also obvious over Queen 1990 and Kurrle.

11. Claims 80 and 81 are Obvious Over Queen 1990 and Kurrle

Claim 80: Claim 80 claims the same "humanized antibody variable domain" as claim 1 (*i.e.*, "comprising non-human CDR amino acid residues which bind an antigen ... and further comprising a Framework Region (FR) amino acid substitution" at residues which completely overlap with claim 1). Like claim 64, claim 80 further recites functional aspects of the humanized antibody, including: (a) noncovalently binds antigen directly; (b) interacts with a CDR; or (c) participates in the V_L-V_H interface ..." Ex. 1003 at ¶¶ 155-58, 186-90, 239-42.

The additional recited elements, which are noted functions of the substituted residues, do not add anything new to the claim. See claim 64, §VI.I.6; Ex. 1003 at ¶240; see also Atlas Powder, 190 F.3d at 1347. Even assuming one could discount the inherency of these functions (which Mylan disagrees with), both Queen 1990 and Kurrle explicitly teach interaction of the framework residues with the CDR as a reason for substitutability. See Ex. 1050 at 16, ll. 4-8; Ex. 1071 at 8, ll. 28-29 and 32-40; Ex. 1003 at ¶¶240-41. For the same reasons as claims 1 and 64 above, see §§VI.I.1 & 6 supra, including the disclosure of framework region substitutions at positions 98L and 36H as provided by Queen 1990 (§§VI.H.1 & I.1 supra), and 4L, **69H**, **73H** and **76H** (§§VI.G.1, G.4 & I.1 *supra*), as provided by Kurrle, as well as the explicit motivation and reasonable expectation of success provided by both Queen 1990 and Kurrle (see §§ VI.G.1 & H.1), claim 80 of the '213 patent is obvious over Queen 1990 and Kurrle.

Claim 81: Claim 81 depends on claim 80, adding "wherein the substituted residue is the residue found at the corresponding location of the non-human antibody from which the non-human CDR amino acid residues are obtained." Both Queen 1990 and Kurrle teach this. See Ex. 1050 at 7, 1. 36-8, 1. 1; Ex. 1071 at 8, 1l. 28-31; Ex. 1003 at ¶242. Claim 81 is also obvious over Queen 1990 and Kurrle.

J. <u>Ground 4</u>: Claim 12 Is Obvious Over Queen 1990 and Kurrle, In View of Furey

The POSITA further would have been motivated to identify residues important for antibody binding, e.g., CDR contact residues and $V_L:V_H$ interaction. Ex. 1003 at \$\\$984-88, 203-204.

Claim 12, which depends on claim 1, recites "wherein the residue at site 66L has been substituted." Furey disclosed the importance of residue 66L as maintaining antigen binding and specificity. *See* Ex. 1053 at Abstract; Ex. 1003 at ¶203-205. Specifically, Furey identified position 66L as interacting with CDR2 of the light chain. Ex. 1053 at Table 4; Ex. 1003 at ¶204.

This directly ties to Queen 1990's and Kurrle's teachings, which provided a POSITA the motivation and reasonable expectation of success to substitute framework region positions that are close enough to interact directly with antigen, as Furey identified with residue **66L**, which a POSITA would have understood as being on a list of substitutable residues in order to maintain antigen binding and specificity. *See* Ex. 1053 at Table 4; Ex. 1003 at ¶204. Claim 12 is thus obvious over Queen 1990

and Kurrle, and further in view of Furey.

K. <u>Ground 5</u>: Claims 73, 74, 77 and 79 are Obvious Over Queen 1990 and Kurrle, In View of Chothia & Lesk and Chothia 1985

Claims 73 and 77: Claims 73 and 77, which both depend on claim 66, recite "wherein the residue at site 78H has been substituted" (claim 73), and "which further comprises amino acid substitutions at sites 71H, 73H and 78H (claim 77). Residue 78H was already known as being important for maintaining antibody conformation, and thus antigen binding and specificity, as identified by Chothia & Lesk and Queen 1990. See Ex. 1062 at Abstract; Ex. 1003 at ¶229. Chothia & Lesk found that "[t]he major determinants of the tertiary structure of the framework are the residues buried within and between the $[V_L \text{ and } V_H]$ domains," including residue **78H** specifically. Ex. 1062 at 903; Table 4; Ex. 1003 at ¶¶230-31. The Background of the '213 patent also recognized the importance of Chothia & Lesk's findings. See Ex. 1001 at col. 3, Il. 1-8 (citing to Chothia, Lesk and colleagues for residues "critically affecting the conformation of particular CDRs and thus their contribution to antigen binding." Citing to Ex. 1062).

Thus, the field, including the '213 patent inventors, already recognized the importance of framework residues, such as **78H**, that are important to maintain antigen binding. Ex. 1003 at ¶230. In view of the importance of **78H**, it would have been obvious for a POSITA to include **78H** as a substitutable residue. Ex. 1003 at ¶231. Claim 73 is obvious over Queen 1990, Kurrle and Chothia & Lesk.

Adding residue **78H** to the combination of residues **71H** and **73H** does not extend patentability. Ex. 1003 at ¶232. These residues were substituted (**71H** and **73H**) in Kurrle, or would have been substituted (**78H**) if necessary. Ex. 1003 at ¶232. Claim **77** is also obvious over Queen 1990, Kurrle and Chothia & Lesk.

Claim 74: Claim 74, which also depends on claim 66, recites "wherein the residue at site 93H has been substituted." Chothia 1985 identified residue 93H as important for maintaining V_L:V_H interactions. See Ex. 1063 at Table 4; Ex. 1003 at ¶225-26. The inventors of the '213 patent also recognized the importance of residues that maintain VL:VH interface contact, as disclosed in Chothia. See Ex. 1001 at 3:1-8, supra; see also Ex. 1050 at 17 (recognizing the importance of "residues essential for inter-chain interactions."). Thus, Kurrle and Queen 1990 provided the explicit motivation as well as reasonable expectation of success to substitute residue 93H for the non-human (e.g., murine) residue, and thus made obvious that residue 93H would have been substituted. Ex. 1003 at ¶225-26. Claim 74 is also obvious over Queen 1990, Kurrle and Chothia & Lesk.

<u>Claim 79</u>: Claim 79 recites "a humanized variant of a non-human parent antibody which binds an antigen, wherein the humanized variant comprises

Complementarity Determining Region (CDR) amino acid residues of the non-human parent antibody incorporated into a human antibody variable domain, and further

comprises Framework Region (FR) substitutions at heavy chain positions 71H, 73H, 78H and 93H, utilizing the numbering system set forth in Kabat."

As above, Kurrle already substituted positions 71H and 73H. *See* Ex. 1071 at Table 6B; Ex. 1003 at ¶234, Padlan Exhibit B. Chothia 1985 disclosed residue 93H as important for maintaining V_L:V_H interactions. Ex. 1063 at Table 4; Ex. 1003 at ¶¶225-26, 235. Finally, Chothia & Lesk disclosed residue 78H as one specifically and independently important for maintaining antigen binding. Ex. 1062 at Table 4; Ex. 1003 at ¶¶230-31. It would have been obvious to a POSITA to combine substitutions at 71H, 73H, 78H and 93H, as taught by Queen 1990, Kurrle, Chothia & Lesk and Chothia 1985. *See* §§VI.G.4, I.1 & I.9, *supra*; Ex. 1003 at ¶¶229-38. Claim 79 is also obvious over Queen 1990 and Kurrle, and further in view of Chothia & Lesk and Chothia 1985.

L. Ground 6: Claims 30, 31 and 33 Are Obvious Over Queen 1990 In View of Hudziak

Claims 30 and 42: Claim 30 of the '213 patent recites "[a]n antibody which binds p185^{HER2} and comprises a humanized antibody variable domain, wherein the humanized antibody variable domain comprises non-human Complementarity Determining Region (CDR) amino acid residues which bind p185^{HER2} incorporated into a human antibody variable domain and further comprises a Framework Region (FR) amino acid substitution at a site selected from the group consisting of: 4L, 38L, 43L, 44L, 46L, 58L, 62L, 65L, 66L, 67L, 68L, 69L, 73L, 85L, 98L, 2H, 4H, 36H,

39H, 43H, 45H, 69H, 70H, 74H, 75H, 76H, 78H and 92H, utilizing the numbering system set forth in Kabat." Claim 42 depends from claim 30, and further recites "wherein the residue at site 66L has been substituted."

Claim 30 is similar to claim 1, differing in the recitation that the CDRs (and antibody) also bind to p185^{HER2}. Claim 30 also includes additional framework sites for substitution at positions 46L, 75H and 76H.

Antibody humanization was developed for a single purpose: realizing the therapeutic promise of monoclonal antibodies for the treatment of human diseases. Ex. 1003 at ¶321; Ex. 1004 at ¶41-45. While monoclonals were capable of targeting antigens (*e.g.*, proteins) in a highly specific manner, immunogenicity issues severely limited the applicability of this technology to human therapeutics. *See* Ex. 1003 at ¶321; Ex. 1004 at ¶42.

A prime molecular target was HER2/*c-erbB-2*, whose amplification in breast cancer patients was correlated with poor prognosis and high relapse rate. *See* Ex. 1021 at Abstract, 1; Ex. 1004 at ¶¶54-75; Ex. 1003 at ¶¶322-23. With respect to the HER2/*c-erbB-2* gene product p185^{HER2}, Hudziak reported that:

- p185^{HER2} was amplified in about 30% of breast cancer tumors; Ex. 1021 at 1; Ex. 1004 at ¶58; Ex. 1003 at ¶322;
- p185^{HER2} "correlated with a negative prognosis and high probability of relapse"; Ex. 1021 at 1; Ex. 1004 at ¶58; Ex. 1003 at ¶322;

- Increased expression of HER-2/*neu* resulted in cellular transformation of the cells and tumorigenesis when the transformed cells were implanted in athymic mice, Ex. 1021 at 1; Ex. 1004 at ¶60; Ex. 1003 at ¶322; and
- High levels of HER-2 gene expression resulted in the cells forming anchorage-independent colonies in soft agar and at low density in low serum concentration, which are characteristics of a transformed phenotype, Ex. 1021 at 1; Ex. 1004 at ¶66; Ex. 1003 at ¶322.

In reviewing Hudziak [Ex. 1021] and other literature, Mylan's expert Dr. Edward T. Ball, a practicing oncologist with clinical experience in developing therapeutic antibodies, concluded the above findings "strongly suggested that the HER-2/*neu* receptor was a ripe target for therapeutic development." Ex. 1004 at ¶61; Ex. 1003 at ¶322-23.

Moreover, a POSITA would have been motivated to develop a monoclonal antibody therapeutic against p185^{HER2}, particularly because of its structural similarity to other growth factor receptors, including epidermal growth factor receptor (EGFR). *See* Ex. 1004 at ¶63; Ex. 1003 at ¶324. This was demonstrated well prior to June 1991 for 4D5, a well-characterized mouse monoclonal antibody targeting p185^{HER2} protein with high affinity, specificity (no cross-reactivity with, for example, EGFR) and efficacy in *in vitro* and *in vivo* studies. *Id.* at ¶65; Ex. 1003 at ¶325. The investigators concluded that 4D5 provided "new potential for diagnostic approaches and

therapeutic strategies for treatment of human malignancies." Ex. 1047 at 4; Ex. 1004 at ¶70; Ex. 1003 at ¶325.

Given published accounts regarding other monoclonal antibody humanization efforts, and the strength of 4D5 as a clinical target, the logical and necessary next step would have been to humanize 4D5. Ex. 1004 at ¶78; Ex. 1003 at ¶325. The 4D5 investigators urged artisans to follow precisely this path:

The muMAb 4D5 also serves as a template for antibody engineering efforts to construct humanized versions more suitable for chronic therapy or other molecules which may be directly cytotoxic for tumor cells overexpressing the HER2 protooncogene.

Ex. 1048 at 10; Ex. 1004 at ¶75 (emphasis added).

Queen 1990 provided the detailed steps for humanizing mouse monoclonal antibodies, such as 4D5, and represented the state of the art of antibody humanization by 1991, teaching humanization of antibody variable domains having non-human CDR amino acid residues that bind to an antigen and are incorporated into a human antibody variable framework domain. Ex. 1003 at ¶118-24, 334. Further, Queen 1990 explicitly disclosed that a POSITA would have had a reasonable expectation that such a humanized antibody would be capable of binding to p185^{HER2}. See Ex. 1050 at Abstract ("the humanized immunoglobulins of the present invention will be substantially non-immunogenic in humans and retain substantially the same affinity as the donor immunoglobulin to the antigen ..."); Ex. 1003 at ¶334. Queen 1990, thus,

provided the explicit motivation to make framework substitutions that would, for example, be more representative of a human residue (Criterion II; *id.* at 15, ll. 22-37), residues that are "immediately adjacent" to CDRs that "likely [] interact with ... the CDR's..." (Criterion III; *id.* at 16, ll. 1-12), and residues that are "in contact", *i.e.*, within about 3 Å of a CDR (Criterion IV; *id.* at 14, l. 14-15, l.2). Ex. 1003 at ¶¶118-24.

Hudziak provided explicit motivation to develop 4D5 for therapeutic use, disclosing "monoclonal antibodies specific for p185^{HER2} (*e.g.*, 4D5) [as] useful therapeutic agents for the treatment of human neoplasias." *See* Ex. 1021 at 7; Ex. 1003 at ¶¶321-25, 333; Ex. 1004 at ¶70. POSITAs would have recognized in June 1991, that 4D5 required humanization before clinical use. *See* Ex. 1048 at 10 ("4D5 also serves as a template for antibody engineering efforts to construct humanized versions more suitable for chronic therapy ..."); Ex. 1003 at ¶¶324-25, 333-34, 337; Ex. 1004 at ¶75. As discussed in §§VI.E.2 & 3, and H.1, *supra*, the particular residues to modify would have included at least **66L**, **98L** and **36H** which likewise appear in claims 30 and 42. Claims 30 and 42 were obvious over Queen 1990 and Hudziak.

Claim 31. Claim 31 recites that "the substituted residue is the residue found at the corresponding location of the non-human antibody from which the non-human CDR amino acid residues are obtained." Queen 1990 explicitly disclosed this

limitation. *See* Ex. 1050 at 5, 1. 36 - 6, 1.1; Ex. 1003 at ¶¶118-24, 335. Claim 31 is also obvious over Queen 1990 and Hudziak.

Claim 33. Claim 33 further adds that "the human antibody variable domain is a consensus human variable domain," which Queen 1990 explicitly discloses. *See* Ex. 1050 at 12, ll. 17-20 ("As acceptor ... use a consensus framework from many human antibodies."); Ex. 1003 at ¶¶118-24, 336. For at least these and the reasons for claim 30, claim 33 is also obvious over Queen 1990 and Hudziak.

M. Ground 7: Claim 42 is Obvious Over Queen 1990 in view of Furey and Hudziak

Claim 42, which depends on claim 30, recites "wherein the residue at site 66L has been substituted." As discussed above, the art taught individual residues to target for humanization for additional reasons; Furey disclosed that residue 66L forms a hydrogen bond contact with CDR2 of the light chain. *See* Ex. 1125 at Table 4; Ex. 1003 at ¶337-39. Following the detailed roadmap of Queen 1990 and Kurrle; recognizing Furey's particular emphasis on 66L to improve binding affinity; and in light of the teachings of Hudziak motivating a POSITA to target p185^{HER2} with an antibody, particularly 4D5, Ex. 1003 at ¶337-39, Ex. 1004 at ¶70, a POSITA would have placed residue 66L on a short list of substitutable residues. Ex. 1003 at ¶337-39. Claim 42 is obvious over Queen 1990 and Kurrle, and further in view of Furey and Hudziak.

N. Ground 8: Claim 60 is Obvious Over Queen 1990 In view of Chothia & Lesk and Hudziak

Claim 60, which also depends on claim 30, recites "wherein the residue at site 78H has been substituted." Chothia & Lesk disclosed a small universe of residues which are "primarily responsible for the main-chain conformations of the hypervariable regions" (*i.e.*, maintaining CDR conformation as Queen 1990 taught), including residue **78H**. *See* Ex. 1062 at Abstract, Table 4; Ex. 1003 at ¶337-39. Following the detailed roadmap of Queen 1990, a POSITA would have looked to Chothia & Lesk and identified FR positions that could interact with or influence CDR conformation, and antigen binding and specificity, including residue **78H**. Ex. 1003 at ¶337-39. Claim 60 is obvious over Queen 1990 and Kurrle, in view of Chothia & Lesk.

O. Secondary Considerations Cannot Preclude Obviousness.

Patent Owner may attempt to assert secondary considerations of nonobviousness, despite no showing of such in the patent. Such evidence would be "insufficient" to "overcome the strong [case] of obviousness" here, *Pfizer, Inc. v. Apotex, Inc.*, 480 F.3d 1348, 1372 (Fed. Cir. 2007). Patent Owner cannot show the required nexus between any purportedly novel feature and any secondary consideration. *See, e.g., Merck & Co. v. Teva Pharms. USA*, 395 F.3d 1364, 1376 (Fed. Cir. 2005); *see also Torrent Pharms. Ltd. v. Novartis AG*, IPR2014-00784 at 12 (PTAB Sep. 24, 2015)("If objective indicia of nonobviousness are due to an element

in the prior art, no nexus exists."). Patent Owner cannot show secondary considerations are commensurate with claim scope given the extraordinary breadth of the challenged claims here. *See, e.g., Ex Parte Takeshi Shimono*, Appeal 2013-003410 (PTAB Apr. 29, 2015). Mylan nonetheless preliminarily addresses potential Patent Owner theories below.

1. The Methods Recited in the '213 Patent Produced No Relevant Unexpected Results.

The '213 patent makes no claim that the claimed methods achieve any unexpected result. To the contrary, the '213 patent recognizes the work of others, including Mylan's expert Dr. Padlan, that residues important for maintaining CDR conformation and binding were well known prior to June 1991. *See* Ex. 1001 at col. 2, 1. 63 – col. 3, 1. 8; Ex. 1003 at ¶¶341-342. Given the extensive prior art, successful antibody humanization was readily achievable, not surprising or unexpected. Ex. 1003 at ¶¶ 341-342; Ex. 1004 at ¶¶46-53.

2. The '213 Patent Satisfied No Long-Felt But Unmet Need.

There was no long-felt but unmet need for humanized mouse monoclonal antibody 4D5. First, the challenged claims' scopes well exceed antibody 4D5 specifically; <u>if</u> 4D5 satisfied any need, the mouse monoclonal antibody 4D5 disclosures, which claimed and disclosed the original mouse monoclonal antibody, satisfied it. *See, e.g.*, U.S. Patent No. 5,677,171 (Ex. 1096); Ex. 1003 at ¶342.

Patent Owner cannot even show the purported invention solved the problem the

specification identified. *See, e.g., Norgren Inc. v. ITC*, 699 F.3d 1317 (Fed. Cir. 2012) (patent obvious where "[prior art patent] solved similar problems in a similar way."); *see also In re PepperBall Techs., Inc.*, 469 F. App'x 878, 882-83 (Fed. Cir. 2012). The '213 patent's purported problem was that "[m]ethods are needed for rationalizing the selection of sites for substitution in preparing [humanized] antibodies" and claimed their invention could provide methods "for the preparation of antibodies that are less antigenic in humans...but have desired antigen binding." Ex. 1001 at col. 3, ll. 53-55 and col. 4, ll. 24-35. Queen 1990, Kurrle and others had already described exactly this process - they set forth why one would desire to humanize and provided a detailed roadmap on exactly how to do it. Any problems identified in the '213 specification had already been solved and explicitly addressed by the prior art. Ex. 1003 at ¶340-42.

3. No nexus/commercial success to Herceptin.

First, none of the heavy chain residues cited in claim 1 are even modified in Herceptin,⁶ and only 1 of the 13 heavy chain residues (78H) cited in claims 30, 62 and 63 is modified in Herceptin. *Second*, the challenged claims are not limited to any

⁶ Mylan presumes that Patent Owner will attempt to rely on Herceptin when presenting its evidence on secondary considerations. Mylan does not concede that Herceptin provides support for any asserted secondary considerations.

particular antibody or even class of antibodies. Ex. 1003 at ¶343. Even claim 30, which recites that the antibody binds p185^{HER2}, is exceptionally broad, not being limited to any specific anti-p185^{HER2} antibody. Therefore, even if Patent Owner can identify one embodiment in its evidence of objective indicia, they will be unable to "demonstrate that untested embodiments falling within the claimed range will behave in the same manner." *Id.* at 4.

Dated: August 2016

Jeffrey W. Guise, Ph.D.

Lead Counsel Reg. No. 34,613

Wilson Sonsini Goodrich & Rosati PC

650 Page Mill Road Palo Alto, CA 94304

Telephone: (858) 350-2225 Facsimile: (650) 493-6811 E-mail: jguise@wsgr.com

CERTIFICATE OF SERVICE

Pursuant to 37 C.F.R. §§ 42.6(e) and 42.105, I certify that I caused to be served a true and correct copy of the foregoing: **PETITION FOR INTER**

PARTES REVIEW OF U.S. PATENT NO. 6,407,213 by Federal Express Next

<u>Business Day Delivery</u> on this day, August 30, 2016 on the Patent Owner's correspondence address of record for the subject patent as follows:

GENENTECH, INC.
1 DNA WAY
SOUTH SAN FRANCISCO, CA
94080-4990

GENENTECH, INC. 460 POINT SAN BRUNO BLVD. SO. SAN FRANCISCO, CA 94080

SIDLEY AUSTIN LLP 2001 ROSS AVENUE Suite 3600 DALLAS, TEXAS 75201

Respectfully submitted,

Dated: August 30, 2016

effrey W. Guise, Ph.D.,

Lead Counsel Reg. No. 34,613

Wilson Sonsini Goodrich & Rosati PC 650 Page Mill Road

Palo Alto, CA 94304

Telephone: (858) 350-2225 Facsimile: (650) 493-6811 E-mail: jguise@wsgr.com