

Sequence analysis

Humanization of antibodies using a machine learning approach on large-scale repertoire data

Claire Marks , Alissa M. Hummer , Mark Chin and Charlotte M. Deane *

Department of Statistics, University of Oxford, Oxford OX1 3LB, UK

*To whom correspondence should be addressed.

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Abstract

Motivation: Monoclonal antibody (mAb) therapeutics are often produced from non-human sources (typically murine), and can therefore generate immunogenic responses in humans. Humanization procedures aim to produce antibody therapeutics that do not elicit an immune response and are safe for human use, without impacting efficacy. Humanization is normally carried out in a largely trial-and-error experimental process. We have built machine learning classifiers that can discriminate between human and non-human antibody variable domain sequences using the large amount of repertoire data now available.

Results: Our classifiers consistently outperform the current best-in-class model for distinguishing human from murine sequences, and our output scores exhibit a negative relationship with the experimental immunogenicity of existing antibody therapeutics. We used our classifiers to develop a novel, computational humanization tool, Hu-mAb, that suggests mutations to an input sequence to reduce its immunogenicity. For a set of therapeutic antibodies with known precursor sequences, the mutations suggested by Hu-mAb show substantial overlap with those deduced experimentally. Hu-mAb is therefore an effective replacement for trial-and-error humanization experiments, producing similar results in a fraction of the time.

Availability and implementation: Hu-mAb (humanness scoring and humanization) is freely available to use at opig.stats.ox.ac.uk/webapps/humab.

Contact: deane@stats.ox.ac.uk

Supplementary information: [Supplementary data](#) are available at *Bioinformatics* online.

1 Introduction

Since the first monoclonal antibody (mAb), muromonab, was approved by the US FDA in 1986, the antibody therapeutics market has grown exponentially, with 6 of the top 10 selling drugs in 2018 being mAbs (Lu *et al.*, 2020). Therapeutic mAbs and antibody-related products such as Fc-fusion proteins, antibody fragments, nanobodies and antibody-drug conjugates are now the predominant class of biopharmaceuticals, representing half the total sales of all biopharmaceutical products (Ecker *et al.*, 2015). These therapeutics treat a range of pathologies including but not limited to cancer, multiple sclerosis, asthma and rheumatoid arthritis. As of September 2020, 93 therapeutic mAbs have been approved by the US FDA and at least 400 others are in development (Raybould *et al.*, 2020).

Many therapeutic antibodies are derived from natural B-cell repertoires of mice, or mice with an engineered human germline repertoire (Lu *et al.*, 2020). However, antibodies developed in animal models are often not tolerated by humans and can elicit an immune response—this property is known as immunogenicity. Immunogenic responses can negatively impact both safety and pharmacokinetic

properties of the therapeutics and can result in the production of neutralizing antibodies that lead to loss of efficacy (Lu *et al.*, 2020). This can pose a significant barrier to the development and approval of therapeutics (Chirino *et al.*, 2004). To combat the immunogenicity of mAbs, various techniques to engineer murine antibodies by substituting part of their sequence with human ones are used. These include chimerization (Morrison *et al.*, 1984) and humanization (Jones *et al.*, 1986). The former involves the combining of a murine variable domain with human constant region domains, and the latter involves grafting the murine complementarity-determining region (CDR) sequences into a human scaffold. Early studies have suggested that more human-like sequences demonstrate lower levels of immunogenicity (Hwang and Foote, 2005). While multiple techniques have been developed to obtain fully human mAbs, humanized antibodies remain the predominant class of mAb making up ~50% of therapeutics in development (Raybould *et al.*, 2020).

The aim of humanization is to reduce immunogenicity while preserving the efficacy of the therapeutic. Typically, human frameworks with high homology to the original sequence of interest are chosen as a scaffold (Safdari *et al.*, 2013). Some murine residues in

framework regions, referred to as Vernier zone residues, affect the conformation of CDR loops and may therefore be retained to preserve antibody affinity. The humanization process remains a bottleneck in mAb development, often based on trial-and-error, involving arbitrary back mutations to restore efficacy or reduce immunogenicity (Williams *et al.*, 2010).

An effective humanization protocol must be systematic and be able to identify the humanness of a sequence with little error. Higher humanness scores should also be linked with lower levels of immunogenicity. Multiple computational methods have been developed to assess antibody humanness. Traditional humanness scores are based on pairwise sequence identity methods between the sample and a set of reference (most often germline) human sequences, for example, a score corresponding to the sequence identity of the closest germline sequence or the average among a set of sequences (Gao *et al.*, 2013). More recent models take account of both preferences of particular residues and pair correlations between amino acids (e.g. Choi *et al.*, 2015; Clavero-Alvarez *et al.*, 2018; Olimpieri *et al.*, 2015; Seeliger, 2013; Wollacott *et al.*, 2019). A multivariate Gaussian model utilized a statistical inference approach (Clavero-Alvarez *et al.*, 2018). This method could distinguish human from murine sequences accurately, but the score demonstrated only a weak negative correlation to experimental immunogenicity levels. More recently, a deep learning approach utilizing a bi-directional long short-term memory (LSTM) model demonstrated best-in-class performance in discriminating between human and murine sequences (Wollacott *et al.*, 2019). However, while making advances, these approaches are subject to limitations including the need for manual input (Olimpieri *et al.*, 2015), the requirement for a solved or modelled antibody structure (Choi *et al.*, 2015), the lack of a humanization protocol (Gao *et al.*, 2013) and limitations in the size of the training dataset (Clavero-Alvarez *et al.*, 2018; Seeliger, 2013; Wollacott *et al.*, 2019).

The recent growth of publicly available antibody sequences has created many opportunities for large-scale data mining. The Observed Antibody Space database (OAS) (Kovaltsuk *et al.*, 2018b) is a database of Ig-seq outputs from 80 studies with nearly 2 billion redundant antibody sequences across diverse immune states and organisms (although primarily human and mouse). OAS is ideal for data mining due to its size, consistent IMGT numbering, and because the sequences represent natural mature antibodies produced *in vivo*.

Utilizing machine learning and the extensive OAS sequence data, we have developed a method that overcomes the limitations of existing techniques. We have constructed random forest (RF) classifiers that accurately distinguish between each human V gene and non-human variable domain sequences. The ‘humanness’ scores produced by our RF classifiers exhibited a negative relationship with observed immunogenicity levels. We used these models to build Hu-mAb, a computational tool that can systematically humanize VH and VL sequences of interest by suggesting mutations that increase humanness. Hu-mAb humanizes the sequence in an optimal manner, minimizing the number of mutations made to the sequence to limit the impact on efficacy. The mutations made by our humanizer were found to be very similar to those made in experimental therapeutic humanization studies that produced sequences with low immunogenicity. Hu-mAb offers a powerful alternative to time-consuming, trial-and-error-based approaches to reducing immunogenicity. Our algorithm is entirely automated, does not require an input structure and uses more sequences for model training than any existing method. Hu-mAb, both humanness scoring and humanization, is freely available at opig.stats.ox.ac.uk/webapps/humab.

2 Materials and methods

2.1 Preparation of OAS antibody sequence datasets

All IgG VH and VL sequences were downloaded from the OAS database (August 2020). Human sequences were split by their V gene type—for example, V1–V7 for VH sequences. Redundant sequences, sequences with cysteine errors (Kovaltsuk *et al.*, 2018a)

and sequences with missing framework 1 residues (residues preceding CDR1) were removed. The total dataset included over 65 million non-redundant sequences (Supplementary Tables S1 and S2). The non-redundant non-human (negative) sequences comprised over 13 million sequences from three species (mouse, rat and rhesus) (Supplementary Fig. S1).

2.2 Training and testing the RF models

All models were trained using the scikit-learn Python module with default parameters unless stated otherwise. RF binary classifiers for each V gene type were trained with their respective set of V gene sequences and the entire set of negative sequences. For example, the VH V1 model was trained on all human VH V1 sequences (labelled as the positive class) and all VH negative sequences (labelled as the negative class). We trained separate classifiers for each human V gene type to achieve a realistic, V gene type-specific sequence representation, rather than a non-physiological representation of mixed V gene types. Eighty percent of the dataset was used for training, 10% for validation and 10% for testing. Performance plateaued after 100–200 estimators and therefore each RF classifier was trained with 200 estimators. The performance of the RF models was assessed by determining their ability to correctly distinguish human sequences of a specific V gene type from those originating from other species. The validation set was utilized to set the classification threshold according to the value that maximizes the Youden’s J statistic (YJS) (calculated as $YJS = \text{sensitivity} + \text{specificity} - 1$). It was found that the threshold that maximizes the YJS was very similar to the threshold that maximizes the Matthews correlation coefficient (Supplementary Table S7). This classification threshold was then used for calculating YJS values of the test set and for classification of therapeutic datasets. In addition, receiver operating characteristic (ROC) curves were generated and area under curve (AUC) scores for each model were calculated in order to assess performance.

2.3 Training and testing the LSTM models

Identical training (excluding negative sequences), validation and test sets were used for the LSTM models. The method to construct the LSTM models followed that described in Wollacott *et al.* (2019). As with the RF models, the validation set was used to set the classification threshold for the test dataset.

2.4 VL Kappa and lambda classifier

An RF model to classify whether a light chain sequence is of type kappa or lambda was trained on 25% of the total human VL dataset (12 million sequences). Testing of the model demonstrated perfect accuracy—it correctly classified every sequence as kappa or lambda within the entire VL dataset (both human and negative).

2.5 Sequence alignments

All antibody sequences were aligned and numbered using the IMGT scheme with the ANARCI software (Dunbar and Deane, 2016).

2.6 Therapeutic antibody dataset

All approved and phases 1–3 antibody therapeutics were obtained from Thera-SABDab (Raybould *et al.*, 2020) and were aligned and IMGT numbered by ANARCI (August 2020). Only mAbs with both a VH and VL sequence were included; this gave a set of 481 therapeutics (Supplementary Material Section 1C). Each therapeutic has an International Nonproprietary Name (INN) assigned by the WHO (Parren *et al.*, 2017). The INN infix preceding the suffix ‘-mab’ is determined by the origin of the therapeutic. Thus, the origin of each therapeutic was obtained from its source infix (Supplementary Fig. S2 and Table S5). Therapeutics named in 2017 onwards no longer followed this nomenclature and their origins were obtained from the IMGT database for therapeutic monoclonal antibodies (IMGT/mab-DB) (Poirion *et al.*, 2010). For 25 therapeutics, we also obtained the precursor sequences prior to their humanization, in addition to their humanized sequences. The Supplementary Material contains a list of all 481

therapeutics and their origin, as well as a list of the 25 experimentally humanized therapeutic sequences and their precursors.

For each of these 481 therapeutics, we calculated the humanness scores of the VH and VL sequences. The VL kappa/lambda classifier (Section 2.4) was used to classify VL sequences as kappa or lambda prior to humanness scoring. A sequence was considered ‘human’ if it achieved a humanness score above the YJS threshold for any V gene. The combined VH/VL sequences were classified as human if both separate VH and VL scores exceeded the respective YJS thresholds.

2.7 Anti-drug antibody response levels of therapeutics

Anti-drug antibody (ADA) responses of patients were obtained for 217 therapeutics from clinical papers using an identical approach to that described in Clavero-Alvarez *et al.* (2018). When multiple ADA levels are reported for the same therapeutic, the mean between the minimal and maximal reported value is used. We then obtained the sequences of 10 additional therapeutics, for which we had ADA response data but which were not included in Thera-SAbDab, from Clavero-Alvarez *et al.* (2018). The complete list of therapeutics together with observed immunogenicity levels can be found in the [Supplementary Material](#).

2.8 Hu-mAb protocol

The input sequence, specific chain type (VH, VL kappa or VL lambda), V gene type and target humanness score were used as inputs. To compare Hu-mAb to experimental mutations, for the therapeutic cases we set the Hu-mAb target score as the humanness score of the experimentally humanized sequence. Every possible single-site mutation within the framework region of the input sequence was made ([Supplementary Fig. S3](#)). This generated a set of mutated sequences which were then scored by the relevant RF model. The humanness scores of the mutated sequences were ranked and the top scoring sequence was selected. This process was repeated with the newly selected sequence until the target humanness score was achieved. We carried out this humanization approach for each of the 25 therapeutics for which we had the precursor and experimentally humanized sequences ([Supplementary Material](#) Section 3G).

To investigate the importance of having separate V gene type-specific classifiers, we conducted a negative control analysis in which we humanized each of these 25 therapeutics sequences using an RF classifier corresponding to a different V gene type than the experimentally humanized sequence. For each therapeutic, we scored the sequence humanness and selected the RF classifier with the lowest humanness score for humanization. In the case where multiple classifiers had the same, lowest humanness score (e.g. a score of 0), the classifier for humanization was selected at random from those with the lowest score.

3 Results

3.1 Classification performance of our RF models on OAS sequences

RF models were generated by training on the OAS IgG dataset (see Section 2). Each model was created as a binary classifier—trained on human antibody sequences (either VH, VL kappa or VL lambda) of a specific V gene type as the positive class and all non-human sequences of the respective chain type as the negative class. Different classifiers were constructed for each V gene as principal component analysis (PCA) demonstrated clear clustering of sequences by their respective V gene type ([Supplementary Fig. S4](#)). The performance of the RF models was assessed by determining their ability to correctly distinguish human sequences of a specific V gene type from those originating from other species. We used the validation set to determine the classification thresholds as the value that maximizes the YJS (see Section 2). Performance on the test set was then calculated using the chosen threshold for each model. Extremely high performance was observed across all models, achieving AUCs (area under the ROC curve) close to 1 or 1 ([Supplementary Table S8](#)). Similar YJS values were also seen in both validation and test sets with all

models scoring ≥ 0.999 . All the VH models perfectly discriminated between human and negative sequences in both validation and test sets. Performance on the light chain was also extremely high, albeit not perfect—this may be due to the greater amount of negative training data available for the VH models (>12 million sequences) compared to that of kappa ($\sim 950\,000$) and lambda ($\sim 650\,000$) models. We also assessed model performance on a subset of our test dataset limited to sequences with $<97\%$ sequence identity with any training/validation sequence [identified using CD-HIT (Fu *et al.*, 2012)] and found no drop off in performance ([Supplementary Table S9](#)).

3.2 Comparison of RF models to previous LSTM models

Recent work has used an LSTM model for predicting humanness (Wollacott *et al.*, 2019). We generated LSTM models with our dataset of sequences (see Section 2) and performance was compared to our RF models. Across all 22 models (each chain and each V gene type), the RF model outperformed the respective LSTM model on both AUC and YJS scores ([Supplementary Table S10](#)). None of the LSTM models was capable of completely discriminating between human and negative sequences. We suspect our RF models produce better results because they are trained using both positive and negative (non-human) data, whereas LSTM models were only trained on positive human sequences.

3.3 Classification of therapeutics

A set of 481 antibody therapeutics (phase I to approved) was obtained from Thera-SAbDab (Raybould *et al.*, 2020) (see Section 2). Each VH and VL sequence was scored by the respective set of RF classifiers (VH, VL kappa or VL lambda) and was classified as human if a single model scored it as human (above the YJS threshold). In the case of VL sequences, we built and used an additional RF model to first discriminate whether the sequence type is kappa or lambda (see Section 2). [Figure 1](#) shows the proportion of therapeutics classified as human (split by origin) by their chain type (VH or VL) and combined (requires both VH and VL to be classified as human). All but 1 of the 176 human sequences were classified as human, and all 14 mouse sequences as non-human. For the one human sequence not classified as such (VH + VL), the light chain humanness score (0.850) fell slightly short of the respective humanness threshold (0.856). Overall our RF models classify more therapeutics as human, as the human content of the antibody sequences increases. This trend is also observed using the LSTM method, but not as clearly—for example, with the LSTM method more of the chi/humanized set are classified as human than the humanized set ([Supplementary Fig. S8](#)). Additionally, more human therapeutics are classified as non-human with the LSTM method.

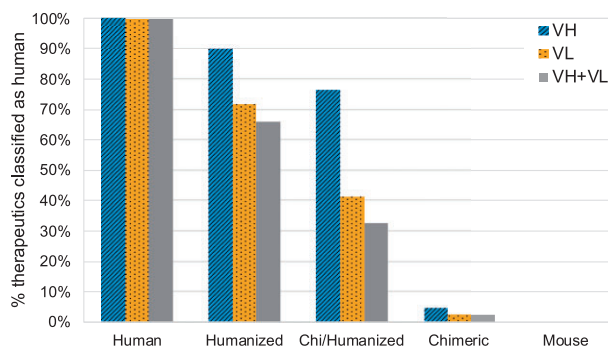


Fig. 1. Percentage of antibody therapeutics classified as human by our RF models, split by their origin: Human (176 sequences), Humanized (214 sequences), Chi/Humanized (34 sequences), Chimeric (43 sequences) and Mouse (14 sequences). Chi/Humanized are sequences which are part humanized and part chimeric. Therapeutics were classified based on their VH and VL sequences separately, as well as combined (to be classified as human, both VH and VL scores had to be above the respective YJS threshold). As the humanness of the therapeutics decreases (left to right), the proportion classified as human also decreases

It might be expected that all chimeric sequences have a completely non-human variable domain as only the constant domains are replaced with human sequence. However, two VH sequences and one VL sequence (out of 43) were labelled as human by our classifiers. This is likely to be because these sequences were of *Macaca irus* origin—a species that was not present in the training dataset from the OAS. Two-thirds of the humanized therapeutics had both VH and VL classified as human. Humanized sequences often have arbitrary back mutations in the framework regions to improve efficacy which might explain why not all humanized sequences are classified as human. Moreover, the INN definition was changed in 2014 such that sequences with a chimeric origin could be given an INN that implied a humanized sequence (Jones et al., 2016). VL sequences had a lower proportion classified as human compared to VH sequences. This could be potentially attributed to the lower number of mutations made in VL sequences (on average 75% of the number of mutations made in VH sequences—see Table 1).

3.4 Relationship of RF model scores with immunogenicity

The aim of humanization is to create a therapeutic that is safe and does not elicit an immune response. A strong predictive score for classification is not sufficient to produce a humanizer as it does not explicitly account for immunogenicity. The relationship of the model scores with observed immunogenic responses, as measured by the appearance of ADAs, was therefore investigated. The fraction of patients with observed immunogenic responses was obtained from FDA labels of approved antibody therapeutics and clinical studies of therapeutics still in clinical trials (outlined in Section 2). There are limitations to this data: for example, there are differences in patient demographic (age, physical conditions, illness), dosage levels and length of dosage of the therapeutic and if the treatment is in combination with other drugs. In addition, the murine therapeutics within the dataset are likely to be inherently biased toward lower levels of immunogenicity as they are approved therapeutics.

We assessed the correlation between the percentage of patients who develop ADAs and the minimum humanness score of a therapeutic's VH and VL chains, as the least human chain is expected to dictate the level of immunogenicity, across 217 therapeutics (Supplementary Fig. S9). We found that higher minimum model scores tend to relate to lower immunogenicity, although the correlation was weak with an R^2 of 0.31. This correlation is substantially higher than the R^2 of 0.18 observed in previous work (Clavero-Alvarez et al., 2018).

We grouped the set of 217 therapeutics by their humanness scores. Figure 2 illustrates this categorization and demonstrates that high humanness scores are linked with low immunogenicity and vice versa. For example, 90% of therapeutics that had both their VH and VL sequence above the YJS threshold exhibited low observed immunogenicity and only one sequence (0.7%) had high immunogenicity. In contrast, <50% of the therapeutics with scores below the YJS threshold had low immunogenicity.

3.5 Hu-mAb: a computational humanizer tool and its application to previously experimentally humanized therapeutics

As high model scores were linked with lower levels of immunogenicity, we used the score to construct a computational humanization tool, Hu-mAb, that suggests optimal mutations that would increase the model score of the input sequence, therefore lowering immunogenicity. Residues in the CDRs are not mutated to maintain antigen-binding properties (described in Section 2). The humanizer should ideally produce as few mutations as possible to reduce efficacy loss of the therapeutic. To investigate the similarity between mutations suggested by Hu-mAb and experimentally derived mutations, experimentally humanized sequences that demonstrated low immunogenicity and for which the precursor sequence was available were collected (Supplementary Tables S3, S11 and S12). The VH and VL sequence of each therapeutic was scored by each RF model, and the

V gene identified by selecting the model that produced the highest score. The precursor sequence was used as the input sequence into the humanizer, along with its target humanness score (the score achieved by the experimentally humanized sequence) and V gene type. An example of the results, for the Campath heavy chain, are shown in Figure 3.

Table 1 compares the mutations made experimentally and those suggested by Hu-mAb for the precursor (unhumanized) sequences of 25 therapeutics. Each of these therapeutics displayed low immunogenicity in their experimentally humanized forms. All precursor sequences were of murine, rat or rabbit origin and most had model scores close to 0 (see Supplementary Material Section 1F/3G for breakdown of immunogenicity and scores). Two therapeutics had precursor sequences scoring above their YJS threshold (VL only for Campath and both VH/VL for Clazakizumab). This is likely due to sequences of their species origin not being present in the training dataset of our models—neither VH/VL rabbit sequences (Clazakizumab) nor VL rat sequences (Campath) were present in the respective training datasets.

Hu-mAb consistently suggested fewer mutations than the number carried out experimentally—on average, Hu-mAb suggested 59% and 58% of the experimental amount for the VH and VL sequences, respectively. Of the mutations suggested by Hu-mAb, an average of 68% and 77% (for VH and VL sequences, respectively) were also made experimentally (overlap ratio, OR). Including mutations to similar residue types (see Supplementary Table S6 for groupings) resulted in an average adjusted OR (AOR) of 77% and 85% for VH and VL, respectively. This shows that the mutations suggested by Hu-mAb are very similar to those made experimentally. In contrast, a randomly humanized sequence would be expected to produce an average OR and AOR of ~2% and ~5%, respectively (see Supplementary Material Section 3H). Hu-mAb is exploiting the information found in the antibody repertoires to more efficiently humanize therapeutic sequences.

We investigated the significance of considering V gene type in humanization by humanizing these therapeutics using an RF classifier of a different V gene type (e.g. humanization of a sequence that is of the VH V1 gene type with the VH V2 classifier). From this, we obtained much poorer humanization results compared to the above. Of the 25 therapeutics, humanization of 19 heavy and 8 light chains was unable to reach the humanness threshold of the experimentally humanized sequence. Where the threshold was reached, an average of 12 and 14 more mutations, for heavy and light chains, respectively, were required to achieve the target humanness score. Furthermore, the OR and AOR, calculated for all mutations suggested even if the threshold was not reached, with the experimentally humanized mutations were on average only 10% and 35% (heavy chain) and 12% and 43% (light chain), respectively. The full results are included in Supplementary Material Section 3I.

3.6 Hu-mAb protocol and RF model analysis

Since experimental humanization procedures often involve grafting of non-human CDRs onto a human framework, it is expected that the framework regions are more important than the hypervariable CDR regions for the classification of human and non-human sequences. Analysis of our RF models' feature importance found that this is true; the key residues for discrimination are mostly found in the framework region (Fig. 4 and Supplementary Figs S5–S7). However, some CDR positions are utilized by the models for discrimination.

Analysis of our Hu-mAb protocol showed that identical mutations (i.e. mutations of position X to residue type Y) do not result in an identical increase in humanness score; the effect depends on the rest of the sequence. Moreover, we found that Hu-mAb occasionally made more than one mutation to the same position in the sequence over the course of the humanization procedure. These observations suggest that our RF models do not consider positions in the sequence independently, but rather they incorporate interactions between residues to more realistically evaluate humanness.

We have also analysed the characteristics of the mutations proposed by Hu-mAb and compared them to those made

Table 1. Comparison between experimental humanization and our computational tool, Hu-mAb

Therapeutic	VH						VL					
	Gene	Unadjusted (%) Overlap ratio	Adjusted (%) Overlap ratio	# Hu-mAb Mutations	# Experimental Mutations	Mutation (%) Ratio	Gene	Unadjusted (%) Overlap ratio	Adjusted (%) Overlap ratio	# Hu-mAb Mutations	# Experimental Mutations	Mutation (%) Ratio
AntiCD28	V3	63	79	19	33	58	KV4	64	73	11	19	58
Campath	V4	75	88	16	39	41	KV1	67	67	3	14	21
Bevacizumab	V3	50	57	14	25	56	KV1	89	100	9	16	56
Herceptin	V3	59	78	27	32	84	KV1	88	88	8	22	36
Omalizumab	V3	62	76	21	34	62	KV1	89	95	19	25	76
Eculizumab	V1	73	73	15	23	65	KV1	83	83	12	20	60
Tocilizumab	V4	64	86	14	23	61	KV1	78	89	9	19	47
Pembrolizumab	V1	73	73	11	23	48	KV3	75	75	12	20	60
Pertuzumab	V3	68	79	19	32	59	KV1	80	90	10	20	50
Ixekizumab	V1	75	75	12	29	41	KV2	78	100	9	12	75
Palivizumab	V2	75	83	12	18	67	KV1	77	92	13	26	50
Certolizumab	V3	61	78	18	31	58	KV1	80	90	10	20	50
Idarucizumab	V4	80	80	15	24	63	KV2	67	67	6	8	75
Reslizumab	V3	50	80	10	21	48	KV1	83	100	6	20	30
Solanezumab	V3	50	70	10	16	63	KV2	88	100	8	10	80
Lorvotuzumab	V3	90	90	10	13	77	KV2	82	82	11	13	85
Pinatuzumab	V3	61	78	23	33	70	KV1	74	79	19	23	83
Etaracizumab	V3	58	83	12	16	75	KV3	62	69	13	25	52
Talacotuzumab	V5	78	83	18	33	55	KV4	73	73	11	16	69
Rovalpituzumab	V1	67	67	21	30	70	KV3	64	79	14	26	54
Clazakizumab	V3	86	86	7	27	26	KV1	75	75	4	22	18
Ligelizumab	V1	64	64	11	21	52	KV3	64	91	11	21	52
Crizanlizumab	V1	64	64	11	23	48	KV1	85	95	20	23	87
Mogamulizumab	V3	67	67	6	15	40	KV2	67	67	6	12	50
Refanezumab	V7	87	87	15	17	88	KV4	92	100	12	17	71
Average		68	77			59		77	85			58
Median		67	78			59		78	88			56

Note: The mutation ratio is the average number of mutations Hu-mAb suggested relative to the number of mutations made experimentally in the framework regions; Hu-mAb never suggests mutations to the CDRs. The overlap ratio is the number of mutations that were both suggested by Hu-mAb and made experimentally, relative to the number of mutations suggested by the Hu-mAb. For the 'unadjusted' overlap ratio, only mutations to identical amino acid types were considered; the 'adjusted' version considers mutations to similar amino acid types to be a match ([Supplementary Table S6](#)).

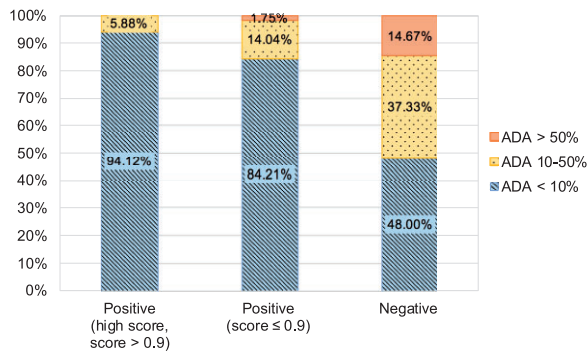


Fig. 2. Relationship between the humanness scores produced by our RF models and experimentally determined immunogenicity. Therapeutics were split into three categories according to the minimum humanness score of the VH and VL chains: positive with a score above 0.9 ['Positive (high score, score > 0.9)'] (85 sequences), above the YJS threshold for the relevant RF model but with a score ≤ 0.9 ['Positive (score ≤ 0.9)'] (57 sequences) and below the YJS threshold ('Negative') (75 sequences). Both the VH and VL sequences have to be above the threshold to be classified as 'Positive'. The immunogenicity of a therapeutic is also represented by three levels: over 50% of patients develop ADAs (orange, solid), 10–50% of patients develop ADAs (yellow, dotted) and under 10% of patients develop ADAs (blue, striped). Therapeutic sequences classified as human by our model tend to have low immunogenicity levels, while sequences classified as not human are more immunogenic

experimentally. In terms of residue types, the mutations proposed by Hu-mAb and experimentally were very similar (Supplementary Fig. S10). Most commonly, mutations were from one hydrophobic residue to another (18% and 20% of all mutations made by Hu-mAb and through experiment, respectively). Least common were mutations involving cysteines (<1% for both Hu-mAb and experiment); importantly the conserved cysteines at IMGT positions 23 and 104 were never mutated, meaning structural viability is maintained (Kovaltsuk et al., 2018a).

The geometry of the antibody binding site is dependent on the orientation of the VH and VL, which is in turn affected by the residues present at the interface between the two domains. The proportion of mutations suggested by Hu-mAb to key VH–VL interface residues is slightly lower than the proportion made by experimental procedures (see Supplementary Table S14), and the OR calculated for these residues is also higher than the average (74%/96% for VH/VL compared to an average across all mutations of 68%/77%). Since Hu-mAb also suggests fewer mutations on average (58–59% of the number made experimentally), the average number of interface mutations per sequence is around half that of experimental

procedures (0.8 vs 1.6 for heavy chains, 0.8 vs 1.8 for light chains). A similar pattern was also observed for the Vernier zone—Hu-mAb proposed fewer mutations to these residues, which are thought to affect CDR conformations (Foote and Winter, 1992) (full details in Supplementary Material Section 3K). This means that the binding properties of the antibody are more likely to be preserved by using Hu-mAb.

4 Discussion

We have developed a novel humanization tool, Hu-mAb, that can humanize both the VH and VL sequences of potential antibody therapeutics. The model is based on RF classifiers that have been trained on large-scale repertoire sequence data and demonstrate very high levels of accuracy in the classification of antibodies by their origin. The humanness scores of the model exhibited a negative relationship with observed experimental immunogenicity. Therefore, sequences that have a higher humanness score are likely to have lower levels of immunogenicity.

Our model is worse at classifying non-human sequences of species that it has not been trained on (as seen with, e.g. the rabbit precursor sequence of Clazakizumab). The non-human sequences within OAS are almost entirely of murine origin, and therefore Hu-mAb is mainly intended for use on murine precursor sequences. We intend to regularly train and update the RF models as new studies of non-human species are added to OAS, potentially widening its uses; however, as most therapeutics of non-human origin are derived from murine sources, our RF models and humanizer Hu-mAb should already be applicable in many cases.

Experimental approaches to humanization are largely a trial-and-error process involving grafting of CDRs onto a completely human scaffold and if efficacy is lost, arbitrary back mutations are made to attempt to restore it (Safdari et al., 2013). Hu-mAb was constructed as a greedy algorithm and is optimized to select the mutations that provide the highest increase in humanness score, thus suggesting as few mutations as possible to reduce the likelihood of impacting the efficacy of the therapeutic. By utilizing RF classifiers that have only trained on a particular V gene type, the humanizer should produce a realistic sequence with a single V gene origin.

Hu-mAb is efficient and only proposes mutations to the key residues in the framework region responsible for humanness; it incrementally suggests additional mutations to reduce immunogenicity if necessary; and back mutations can be suggested in a sequential and non-arbitrary manner (the mutation with the lowest impact on the humanness score). Compared to experimentally humanized therapeutics, Hu-mAb suggested ~60% of the number of mutations, with a high similarity to those suggested experimentally (average AOR of

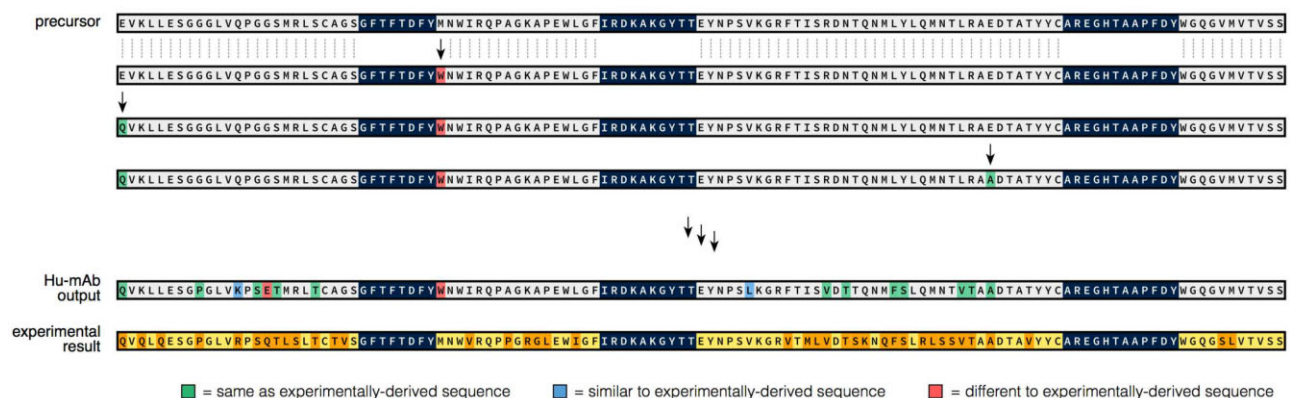


Fig. 3. The Hu-mAb humanization procedure demonstrated using the heavy chain sequence of the therapeutic Campath. The humanized sequence produced experimentally is shown at the bottom of the figure (conserved residues in yellow, mutated residues in orange). Starting with the unhumanized precursor sequence (top), Hu-mAb makes every possible mutation to the framework residues (grey) and selects the one that produces the largest increase in humanness score. CDR residues (dark blue) are not mutated to preserve binding. This procedure is performed iteratively until the humanness score reaches a given threshold. Mutations suggested by Hu-mAb are coloured depending on whether they are the same (green), similar (blue) or different (red) to mutations made experimentally. In this case, Hu-mAb suggested 16 mutations (compared to 39 from the experiment), of which 14 were the same or similar to those derived experimentally

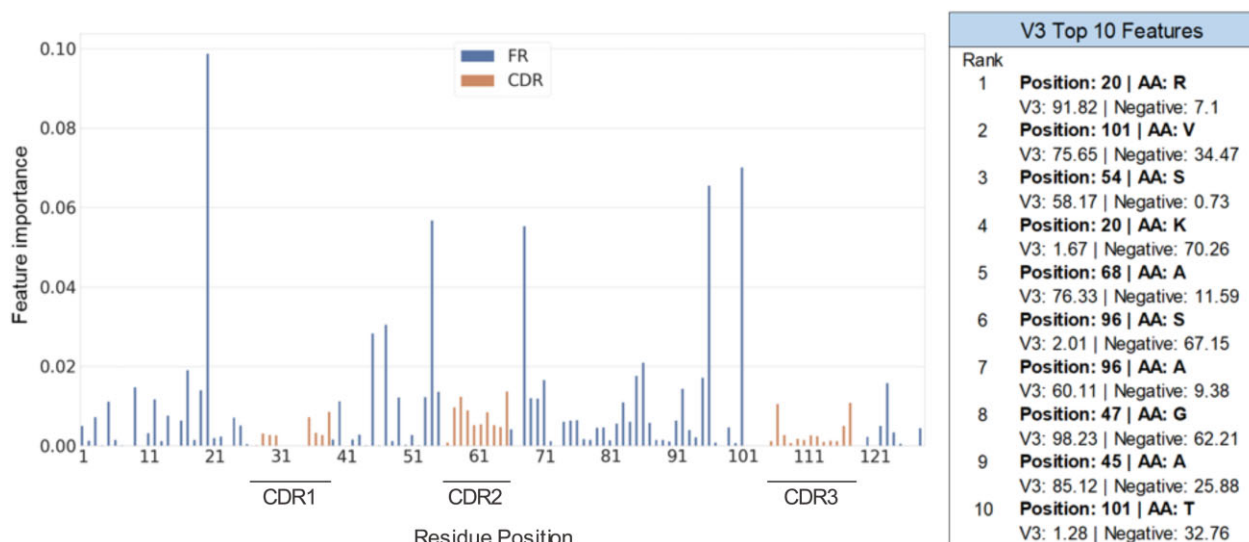


Fig. 4. Feature importance of the VH V3 RF model and its top 10 features. The x-axis consists of the residue positions in a sequential manner (left to right, IMGT numbering scheme). The inset table shows the top 10 features and the percentage frequency of the relevant amino acid type seen within the respective sets of sequences (V3 and negative, or non-human). The most important features likely determine the humanness of the sequence and are mainly located in the framework regions

77/85%). Hu-mAb offers a promising alternative to experimental humanization approaches, allowing mutations to be made in a more systematic and efficient manner, and achieving similar results in a fraction of the time.

There are, however, still remaining challenges in antibody humanization, most notably the humanization of alternative antibody formats (e.g. nanobodies and asymmetric antibodies). The ability to score and increase the humanness of these sequences will improve with further data and investigation of the relationship between immunogenicity, sequence and antibody format.

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