

**Heidelberg University
Institute of Computer Science**

**Project report for the lecture
Advanced Machine Learning**

**Prediction of the next SARS-CoV-2 mutations
by a Transformer based GAN Framework**

<https://github.com/nilskre/AML-covid-project>

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Member contributions

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Came up with the project idea and was the domain expert for the project's biomedical domain. Initial research for the topic and feasibility analysis. Extensive research in the areas of the project's overall composition, data sources, dataset creation and datset insights. Responsible for the data selection and download, the dataset creation, the generation of dataset insights and the pretraining and evaluation of the Transformer model.

Felix Hausberger

Extensive research in the area of ML models, especially the connection between NLP methods and our project, previous work on mutation prediction and different ML model architectures such as Sequence to Sequence, Transformer and GANs. Responsible for the data preprocessing, the model architecture and implementation, the GAN training and evaluation.

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List of Abbreviations

BLEU	Bilingual evaluation understudy
DNA	Deoxyribonucleic Acid
GAN	Generative Adversarial Network
GISAID	Global Initiative on Sharing All Influenza Data
GPU	Graphics Processing Unit
LSTM	Long Short-Term Memory
ML	Machine Learning
NLP	Natural language processing
ORF	Open reading frames
ReLU	Rectified Linear Unit
RKI	Robert Koch Institut
RNA	Ribonucleic Acid
RNN	Recurrent Neural Network
SARS-CoV-2	severe acute respiratory syndrome coronavirus 2
UTR	Untranslated region

0 Project Setup

For a detailed description of how to set up the project, please have a look at <https://github.com/nilskre/AML-covid-project/blob/main/README.md#setup>.

1 Introduction

During the genome replication random mutations can appear. As a consequence the encoded protein sequence could be changed, which can lead to different behavior. If this change increases the fitness, it is probably passed on to the next generation. [6]

A currently well known example for a mutating virus is SARS-CoV-2. Due to the developed vaccines the hope for an end of the pandemic arises. Nevertheless this is only true, if the vaccines, which are developed against the wild type of SARS-CoV-2, also remain effective against new mutations. To enable fast responses to new arising mutations it would be helpful to know the possible next mutations in advance. This can influence the treatment and prevention of diseases, by enabling the development of countermeasures and preventive measures in advance. [6]

Machine Learning (ML), especially Deep Learning enabled improvements in lots of different domains. This work applies Deep Learning to the area of virus genome mutation prediction. Due to the fact, that genome sequences could be treated as text data, methods from the Natural language processing (NLP) area can be applied. The success of Deep Learning for NLP tasks has already been shown in various areas such as text generation, text summarization or translation. [6]

Our research question is whether a ML model can be trained to predict the next possible SARS-CoV-2 mutations. In this paper, we propose three novelties:

- **Model architecture:** A new Generative Adversarial Network (GAN) based architecture influenced by [6]. Our novelty is the usage of transformers instead of Long Short-Term Memory (LSTM) in the seq2seq model.
- **Dataset:** Generation of a dataset for SARS-CoV-2, consisting of 9199 parent child data instances.
- **Application domain:** The training of the network for SARS-CoV-2.

2 Fundamentals and Related Work

2.1 Basics from molecular biology

This chapter gives a brief introduction into the underlying biological domain.

2.1.1 Genome sequences

The human genome is encoded as DNA.

The DNA carries genetic instructions how DNA based organisms develop and behave. It is structured as a double helix and consists of two nucleotide pair combinations: adenine (A), thymine (T) and cytosine (C), guanine (G). [14, p. 8]

2.1.2 Protein biosynthesis

The protein biosynthesis (also known as the central dogma of molecular biology) describes how proteins are produced based on DNA sequences. The proteins then produce characteristics (e.g. human hair color). So the DNA directly influences which human characteristics are developed. [22, p. 6]

The process of the protein biosynthesis can be seen in figure 2. The single steps are described in the following:

1. Transcription: The DNA is transcribed by the RNA-Polymerase into pre-messenger RNA. As part of this process the nucleotide thymine is replaced by uracil (U). [22, p. 9]
2. mRNA Processing: The pre-messenger RNA is transformed into the messenger RNA by removing the non-coding regions (introns). So the main difference between DNA and Ribonucleic Acid (RNA) is that the DNA is structured as a double helix, whereas the RNA consists of a single strain. [22, p. 9]
3. Translation: Ribosomes translate the mRNA into amino acids. Three nucleotides (one codon) decode one amino acid. The matching which codons decode for which amino acid can be seen in figure 3. [22, p. 9]

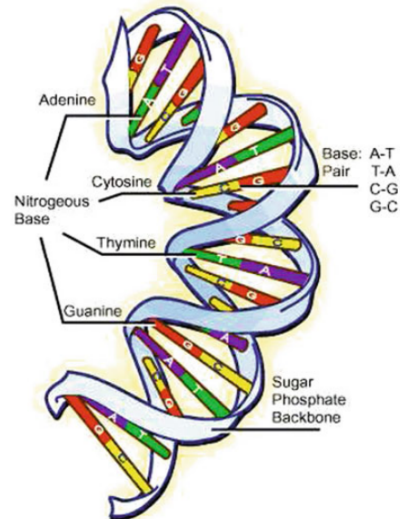


Figure 1: Deoxyribonucleic Acid (DNA) double helix [14, p. 8]

Central dogma of molecular biology

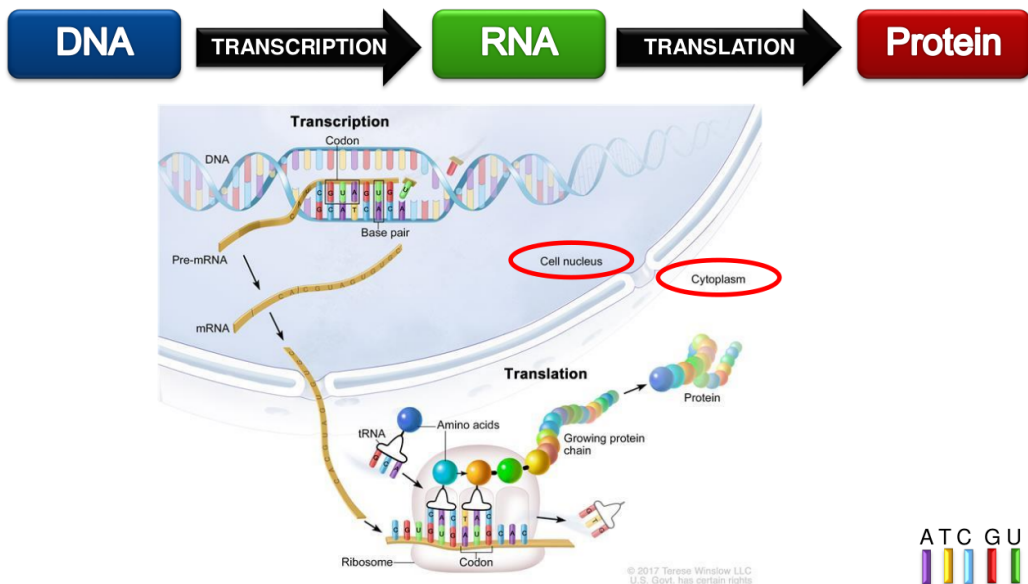


Figure 2: Protein biosynthesis [22]

	U	C	A	G	
U	UUU Phe UUC UUA Leu UUG	UCU UCC Ser UCA UCG	UAU Tyr UAC UAA STOP UAG STOP	UGU Cys UGC UGA STOP UGG Trp	U C A G
C	CUU CUC Leu CUA CUG	CCU CCC Pro CCA CCG	CAU His CAC CAA Gln CAG	CGU CGC Arg CGA CGG	U C A G
A	AUU Ile AUC AUA	ACU ACC ACA ACG	AAU Asn AAC AAA Lys AAG	AGU Ser AGC AGA Arg AGG	U C A G
G	GUU GUC Val GUA GUG	GCU GCC GCA Ala GCG	GAU Asp GAC GAA Glu GAG	GGU GGC GGA Gly GGG	U C A G

Figure 3: Genetic code [22]

2.1.3 Structure of SARS-CoV-2

RNA based viruses (like SARS-CoV-2) use RNA to encode their genetic information. That means that the genetic information is present as a single strained RNA sequence. This sequence contains for example the information which proteins form the surface structure of the SARS-CoV-2 virus and how this surface looks like. [19]

Based on Naqvi et al. [19] the four proteins forming SARS-CoV-2 are:

- spike (S)
- envelope (E)
- membrane (M)
- nucleocapsid (N)

A structural representation of SARS-CoV-2 and a host cell is visualized in figure 4. [19]

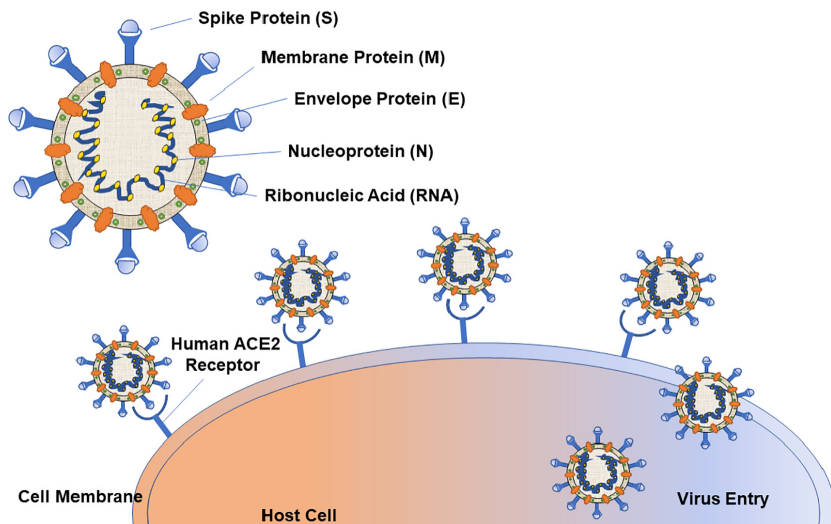


Figure 4: SARS-CoV-2 structure [19]

Furthermore figure 4 shows the RNA in the center. The SARS-CoV-2 RNA consists of about 30,000 nucleotides, which belong to different subgroups, as shown in figure 5. Padded by a 5' Untranslated region (UTR) at the beginning and a 3' UTR in the end, the middle part contains 12 functional Open reading frames (ORF), which contain the four coding regions for the proteins S, E, M and N.

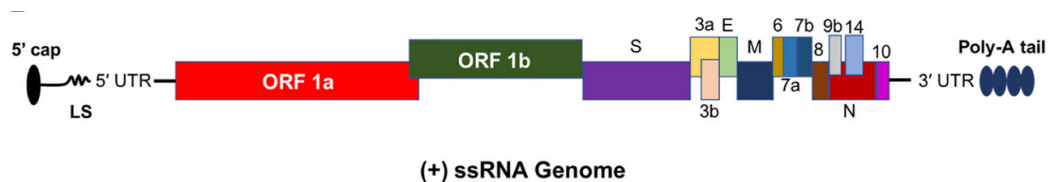


Figure 5: SARS-CoV-2 genome structure [19]

2.1.4 Relationships between biological objects

One main question in biology is to determine relationships between objects, such as individuals or genome sequences. The observed objects are called taxa. Most commonly used for determining these relationships is the phylogenetic analysis which generates a phylogenetic tree. In this phylogenetic tree each leaf node corresponds to exactly one taxa. The inner nodes of the tree are the inferred hypothetical ancestors, which are no taxas. The relatedness between different taxas can be evaluated by their distance. [7]

Figure 6 shows an example phylogenetic tree calculated based on the species genomes. On the right side each leaf node (taxa) corresponds to one current species. From left to right one can see the inferred historical development from the ancestors to the current species. One can see, that Gorillas and Orangutans have earlier developed apart than the other species. Furthermore one can see, that the Chimpanzees and Bonobos share a most recent common ancestor (inner node next to both). That is why they are more related to each other. [15]

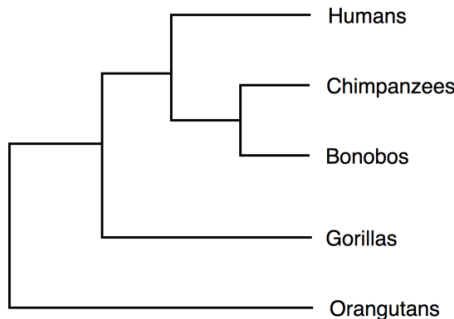


Figure 6: Example phylogenetic tree [15]

2.2 From Probabilistic Language Models to modeling Evolution Theory

A probabilistic language model tries to approximate the probability distribution

$$P(w_1, \dots, w_n) = \prod_{t=1}^n P(w_t | w_1, \dots, w_{t-1}) \quad (1)$$

with w_t being a word at position (time step) t in a sentence of length n . To build language models Recurrent Neural Networks (RNNs) were used to model such probability distributions.

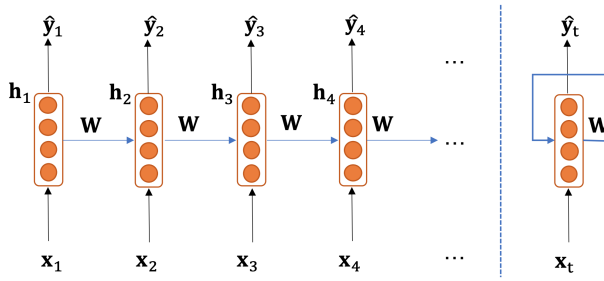


Figure 7: Architecture of a conventional RNN [10]

At each time step t it outputs a probability distribution $P(w_t | w_1, \dots, w_{t-1})$ given the words read so far in the current instance (see Figure 7). Words are read as a vectorized numerical representation, often given by pretrained so-called word embeddings x_t which are lower dimensional and more semantically-enriched compared to simple one-hot encodings. One then calculates the hidden state h_t by

$$h_t = f(W^{(h)}h_{t-1} + W^{(x)}x_t + b_1) \quad (2)$$

and the corresponding output probability distribution by

$$\hat{y}_t = \text{softmax}(U^{(h)}h_t + b_2). \quad (3)$$

The applied weight matrix is always the same for each time step t giving the RNN its name. One can therefore simplify the unrolled RNN architecture on the left side of Figure 7 to the one on the right, where the hidden state is continuously passed as an input to the next time step. To achieve a better convergence behavior during training, one can also provide the expected hidden state of time step $t - 1$ instead of using the predicted hidden state, which is called teacher forcing. RNNs are able to process input of arbitrary length and are by their recurrent character capable to use information from previous time steps. Unfortunately, they are vulnerable to vanishing and

exploding gradient problems. LSTM is a special RNN architecture that solves such vulnerabilities by owning a separate long-term cell state besides a short-term hidden state and is introduced in subsection 2.6. It is able to preserve information over many time steps. [10]

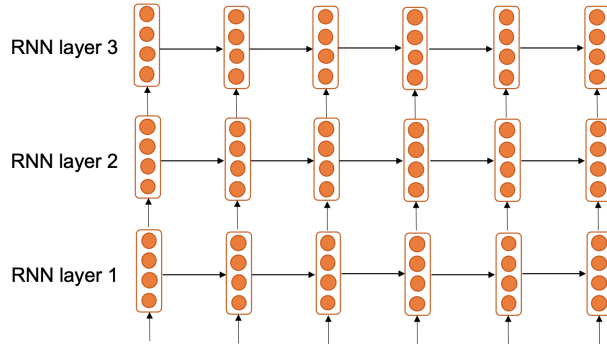


Figure 8: Architecture of a multi-layer RNN [10]

One can also use two RNNs, one traversing a sentence from left to right and another one vice versa, with two different weight matrices to model the probability distribution bidirectionally. One therefore simply concatenates the hidden states of each RNN before applying the weight matrix U and the $\text{softmax}()$ function. Also multi-layer RNN can be utilized to generate higher-order features (hidden states) for the prediction task (see Figure 8). [10]

The RNNs or even better LSTMs architectures used for probabilistic language modeling can be reused in a more complex domain called sequence to sequence modeling for neural machine translation from one language to another. Here one first tries to learn a fixed-dimensional input representation from an input sequence using an encoder architecture based on an LSTM. The so-called context vector is then decoded by a second LSTM into a new sequence of words preserving the grammar but owning a different meaning. [25]

Sequence to sequence models are introduced together with the LSTM architecture in subsection 2.6. Here the connection to evolution theory can be drawn. RNA sequences made of a concatenation of nucleotides¹ can be represented textually using the FASTA format. A sequence to sequence model can then transferably be applied in the domain of RNA sequences to model how RNA-based viruses change their structure to avoid the detection

¹We restrict the representation of nucleotides solely to their nucleobases parts consisting of the distinct nucleobases guanine, adenine, cytosine and thymine. We therefore do not include the phosphate group and the five-carbon sugar components.

by the human immune system but still to preserve their infectivity and evolutionary fitness [12].

2.3 GISAID EpiFlu Data Platform

The data needed for this project consists of the SARS-CoV-2 genome data and the corresponding metadata. The most complete database is hosted by the GISAID initiative [2]. It is a public-private-partnership between the initiative itself and the governments of Germany (official host of the platform), Singapore and the United States of America. [1]

The main aim of GISAID is to enable the fast sharing of epidemic and pandemic virus data. In comparison to other databases such as GenBank the genome sequences are shared fast through GISAID. For enabling this fast sharing the GISAID database is only usable after registration and the redistribution of the data is forbidden, whereas the GenBank database is publicly available. [23]

This leads to the fact, that until september over three million SARS-CoV-2 genome sequences are available through GISAID, whereas only about one million SARS-CoV-2 genome sequences are available through GenBank. Figure 9 shows the selection view of the GISAID database. [1, 16]

The screenshot shows the GISAID database search interface. At the top, there's a navigation bar with links for Registered Users, EpiFlu™, EpiCoV™ (which is highlighted in blue), EpiRSV™, and My profile. Below the navigation bar is a toolbar with icons for EpiCoV™, Search, Downloads, and Upload. The main area is titled "Search" and contains a form with fields for Accession ID, Virus name, Location, Host, Collection, Submission, Clade, Lineage, Substitutions, Variants, and filters for complete, high coverage, low coverage excl, w/Patient status, and collection date compl. A "Reset" button and a "Fulltext" link are also present. Below the form is a table listing 3,364,537 viruses. The columns include: a checkbox column, Virus name, Passage date, Accession ID, Collection date, Submission date, Length, Host, Location, and Originating location. The table lists various entries from Mexico, such as hCoV-19/Mexico/ZAC_IBT_IMSS_2588/2021, hCoV-19/Mexico/ZAC_IBT_IMSS_2587/2021, and hCoV-19/Mexico/SLP_IBT_IMSS_2584/2021, among others. At the bottom of the table, there's a note about accepting the Database Access Agreement and a "Select" button. The footer contains a copyright notice (© 2008 - 2021) and links for Terms of Use, Privacy Notice, and Contact.

Figure 9: GISAID database

2.4 Domain-Specific Methodologies to create Evolutionary Datasets for Mutation Prediction

From databases such as GISAID or GenBank an unordered list of genome sequences and metadata can be downloaded. For training a ML model to detect changes and predict future mutations the dataset must contain parent child pairs.

Berman et al. [6] have done this based on two steps. First they created a phylogenetic tree from the genome sequences. Therefore first all genome sequences are aligned. The initial phylogenetic tree is calculated by Fasttree using the approximate maximum likelihood method. In the next step the tree is refined by a generalized time-reversible (GTR) model and a gamma model. The final phylogenetic tree is achieved by optimizing the branch length and the used models for tree generation. Like in standard phylogenetic trees each leaf node corresponds to one data instance, whereas the inner nodes are inferred from the other data instances. Secondly based on the rooted phylogenetic tree, parent-child pairs are calculated. Therefore first a marginal ancestral sequence reconstruction is executed based on the phylogenetic tree from the previous step. Now also the inner nodes correspond to data instances. In the next step BioPython's [8] Bio.Phylo package is used to determine parent-child pairs. From each edge one parent-child pair is generated. [6]

Mohamed et al. [17] used existing datasets from Ogali et al. [20] in their work about mutation prediction. Ogali et al. [20] performed a phylogenetic analysis. After selecting only genome sequences with high quality, they removed duplicate sequences and added reference sequences to the dataset. Now the alignment is executed. In the next step MEGA (Molecular Evolutionary Genetics Analysis) is used to create the phylogenetic tree. This is done by using the maximum likelihood method, the best-fit general time-reversible (GTR) model and gamma-distributed rate variation.

Hadfield et al. [11] have created the nextstrain project. It aims to examine pathogen's spread and evolution by providing a viral genome database, a bioinformatics pipeline for phylodynamics analysis and a visualization platform. Steps executed by the bioinformatics pipeline for phylodynamic analysis are: "subsampling, alignment, phylogenetic inference, temporal dating of ancestral nodes and discrete trait geographic reconstruction, including inference of the most likely transmission events" [11]. For calculating the phylogenetic tree, TreeTime's maximum likelihood method is used. [11]

2.5 Previous Work on Mutation Prediction

Even before the rise of Covid-19 there had been studies trying to predict mutations of RNA viruses. In the collection of [27, 28, 29] the authors predict the mutation positions in hemagglutinins from influenza A virus using logistic regression and plain neural networks and then use the resulting amino acid mutating probabilities to derive possible mutated amino acids. The same approach is further used for H5N1 neuraminidase proteins.

Salama et al. [21] proved that nucleotides in an RNA sequence can change based on their local neighborhood. Neural networks are used to predict new strains of the Newcastle virus and subsequently a rough set theory based algorithm is introduced to extract the according point mutation patterns.

Mohamed et al. [17] used a more modern sequence to sequence approach based on LSTMs to learn nucleotide mutations between time-series species of H1N1 Influenza virus and the Newcastle virus as mutations can also be influenced by long-distance relations of amino acids. Therefore one hot-encoded RNA sequences of a parent generation preprocessed to words is given as an input and the output is the predicted offspring generation evaluated by accuracy to the compared true offspring generation. The achieved accuracy in this paper is questionably high with 98.9% on the H1N1 Influenza virus and 96.9% on the Newcastle virus, possibly because of overfitting to the few 4.609 samples for H1N1 Influenza virus and only 83 for the Newcastle virus. Our approach therefore tries to increase the number of samples available for training when building the dataset.

Our approach will neither use any of the just mentioned architectures, but uses a transformer based architecture coupled with a GAN-style training architecture. Nevertheless a short introduction into sequence to sequence models and the underlying long short-term memory components shall be given to better point out our architectural decisions.

2.6 Sequence to Sequence Models based on Long Short-Term Memory

The original LSTM unit was introduced by Hochreiter and Schmidhuber in [13] and can be used for language modeling instead of using plain RNNs to prevent running into vanishing or exploding gradient problems [24]. The architecture of an LSTM is shown in figure 10.

It consists of a hidden state h_t and an additional cell state c_t . The cell state stores long-term information and is used to derive a new hidden state. Information flows through three different gates inside the LSTM. The forget gate is used to control which parts of the cell state are potentially carried

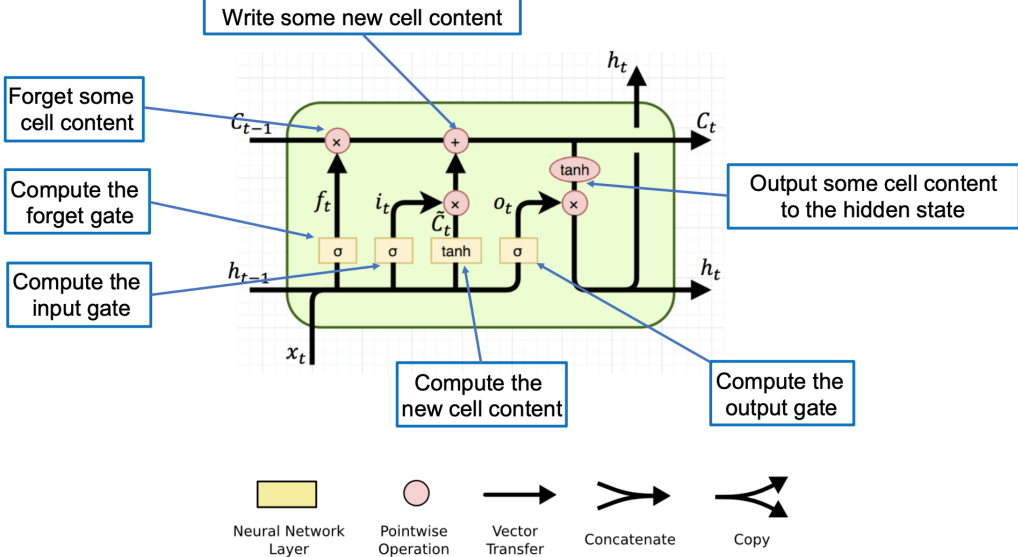


Figure 10: Architecture of an LSTM [10]

on to the next time step, the input gate is responsible to decide which parts of the cell state should be updated and the output gate determines what is being passed on as the new hidden state. All three gates depend on the previous hidden state and the current input. They provide factors limited to the interval $[0,1]$ by the sigmoid function and are multiplied with the cell state, the changes to be added to the cell state and the new hidden state derived from the cell state. Through the cell state an LSTM therefore makes it possible to capture long-distance dependencies. [10]

Sutskever et al. [25] introduced sequence to sequence learning following a multi-layer encoder-decoder style model architecture. One layer consists of one LSTM that is used as an encoder to learn a large fixed-dimensional vector representation of a size-unrestricted input sequence called the context vector. This vector consists of the last cell and hidden state of the encoder and incorporates the structure of the input sequence helping the following decoder LSTM to provide qualitative predictions for the output sequence. The second LSTM therefore serves as a beam search² decoder to map the context vector to a corresponding output sequence whose length does not need to match with the length of the input sequence. The output probability

²Do not choose the most probable word but the B most likely word hypothesis and pass them to the next time step in the LSTM. Whichever hypothesis results in the lowest loss is kept. To avoid combinatorial explosion limit the beam size.

distribution is therefore given by the equation

$$p(y_1, \dots, y_{T'} | x_1, \dots, x_T) = \prod_{t=1}^{T'} p(y_t | v, y_1, \dots, y_{t-1}) \quad (4)$$

with v being the context vector. Using an LSTM is preferred over a normal RNN as it is used to capture the long range temporal dependencies of the input data. The encoder-decoder architecture uses four layers in total partitioned onto four Graphics Processing Units (GPUs). A corpus of 160k words for the input sequence and another one of 80k words for the target sequence was used to create the word embeddings of dimension 1000. Unknown words were replaced by a *UNK* token. The sequence to sequence model approach was evaluated for neural machine translation and reached a 34.81 BLEU score. One finding during training was that reversing the input sequence introduces many short term dependencies as the minimal time lag of the problem is reduced making optimization easier. [25]

2.7 Applying Generative Adversarial Networks

Using a plain sequence to sequence model for mutation prediction does not necessarily guarantee that the generated sequences are evolutionary offsprings of a parent generation as not being included as is in the ground truth data. The generated sequences might occur realistic and biologically relevant, but a plain sequence to sequence architecture does not inherently check for natural parental descent and therefore does not make sure whether the predicted mutations lead to improved fitness. Berman et al. [6] developed a novel sequence to sequence framework based on the GAN idea to predict genetic mutations and future biological populations of the influenza virus (see Figure 11). MutaGAN describes a sequence to sequence generator within an adversarial framework that predicts protein sequences augmented with possible mutations. By using a sequence to sequence generator and a discriminator specialized on separating fake evolutionary mutations from real ones, one can then guarantee to a certain degree that the evolutionary parent-childhood coherence is given. In MutaGAN a mutation is considered correct if the change in amino acid and location within the RNA sequence is equal to the parent's true offspring. [6]

MutaGAN's generator consists of a sequence to sequence model. The encoder is built from of a bidirectional LSTM and the resulting context vector of dimensionality 512 is combined with noise from a standard normal distribution. The decoder LSTM predicts the resulting sequence of amino acids greedily using the *argmax()* of the *softmax()* output for every time step. The discriminator is trained on three different parent-child pair configurations

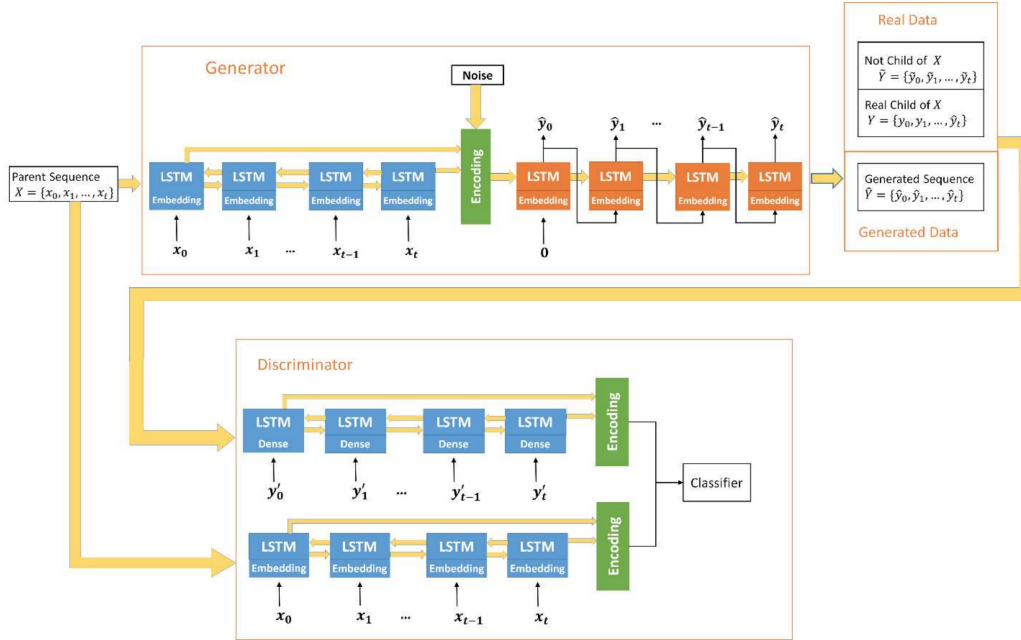


Figure 11: Architecture of MutaGAN [6]

to optimally compete against the generator, real parent-child pairs, real parent and generated child pairs and pairs of real sequences that are not parent-child pairs. Two bidirectional LSTM encoders sharing the same weight matrix as the encoder of the generator produce a fixed-dimensional encoding of the provided parent-child pair used for classification of evolutionary descent by a plain neural network. The child encoder in the discriminator uses a dense layer having the same weight matrix as the embedding layer of the encoder of the generator. The dense layer is required to directly input the predicted child sequence as its probability distribution rather than the final output after applying the *argmax()* as it would not enable backpropagation to train the generator. In case of a true child sequence is given a simple one-hot encoding is used. [6]

Interestingly Berman et al. [6] state planned improvements by utilizing bigger datasets acquired from the GISAID database EpiFlu and by using more length-robust attention-based models to directly work on nucleotide sequences instead of sequences of amino acids. Therefore transformers and the attention mechanism are introduced in the following section.

2.8 Transformer and Attention Mechanism

Using a sequence to sequence model as introduced based on an encoder-decoder architecture of LSTM cells has some drawbacks. First the input needs to be process sequentially in time steps which makes parallelization difficult and increases training time, especially for longer sequences. Furthermore the hidden and cell state vector passed through every timestep tries to encode information of *all* previous time steps without knowing which information is especially important for the current time step. This not only makes long distance dependencies hard to capture, but also might not get the prioritization of the previous time step inputs right. [5, 26]

To tackle these problems the so-called transformer architecture was introduced in [26]. It feeds an entire sequence into the encoder to be processed in parallel denying any concept of recurrence or convolution. Only using a so called self-attention mechanism the transformer makes sure that during processing every input position, each of them receives the information that is most important to them. This way modeling long dependencies becomes much more easy compared to LSTMs as the view on the input sequence is more global. In this architecture, the encoder also passes all computed hidden states of every position to the decoder, which therefore [26] can generate the target sequences based on more semantically enriched features and also in parallel for every position. [26]

The transformer architecture achieved state-of-the-art quality results with a BLEU score of 41.8 on the WMT 2014 English-to-French translation task (cf. BLEU score of [25] was 34.8), while still being much faster to train due to its parallelization capabilities. State-of-the-art results are already achieved after just twelve hours of training on eight P100 GPUs. As this is still far beyond the scope and resources given for this project, this projects provides a proof-of-concept transformer model trained for far shorter and fewer sequences as one would need to predict entirely new RNA sequences. [26]

First the transformer architecture should be introduced in the following:

The transformer consists of an encoder and a decoder part just as normal sequence to sequence models. One layer of the encoder contains a self-attention layer before passing the hidden state representations of an input sequence to a feed forward neural network consisting of two linear transformations separated by Rectified Linear Unit (ReLU). Thus it makes sure that semantic and contextual dependencies between different positions in the input sequence are also modeled. Also skip connections are added for both components with subsequent layer normalization. [4, 26]

To calculate the self-attention output of a sequence, a query Q , key K

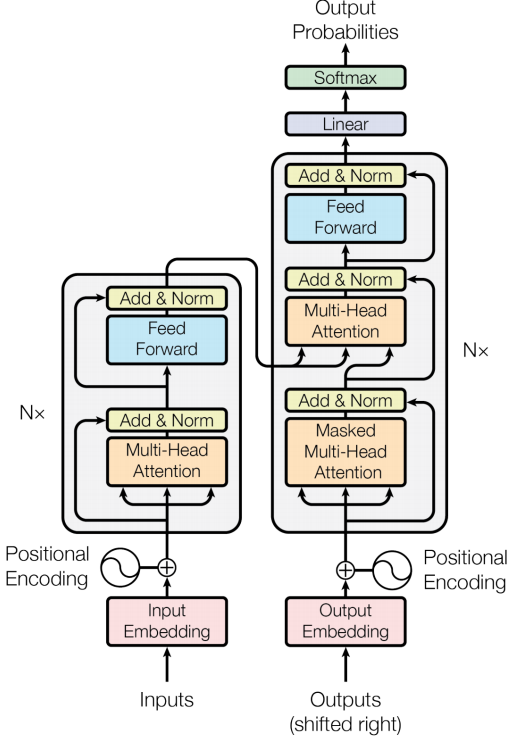


Figure 12: Architecture of the transformer [26]

and value V matrix is calculated through multiplication of the sequence representation³ with learned transformation matrices. Each row of the three received matrices corresponds to a specific input position. The output of the self-attention layer is then calculated through

$$(ScaledDot - Product)Attention(Q, K, V) = softmax\left(\frac{QK^T}{\sqrt{d_k}}\right)V. \quad (5)$$

The dot product between Q and K calculates scores that are used as weighting factors for V . This models for every input position how relevant all other input positions are for each of the hidden state encodings. The scores are divided by the square root of the dimensionality of the query/key values of the hidden state representations, which is chosen to be 64 (square root eight), to receive more stable non-vanishing gradients. The softmax guarantees that the positional scores lie between zero and one and add up to one. Note that the score of the current input position itself will most likely have the highest score. [4, 26]

³A matrix containing the hidden state representations of every input position

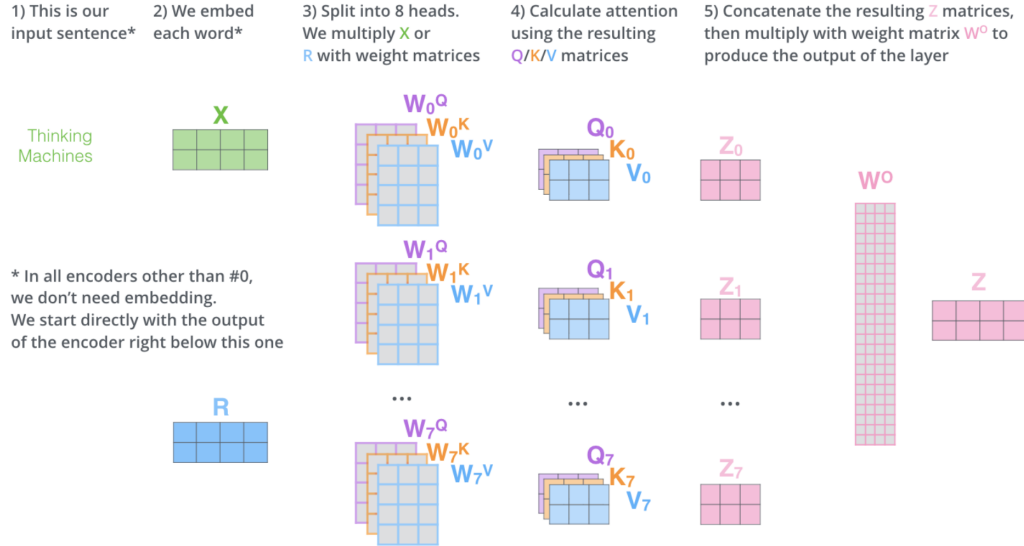


Figure 13: Multi-head attention architecture [4]

One can expand self-attention towards a multi-head attention. In this configuration each head produces its own query, key and value matrices and therefore its own self-attention output in its own representation subspace. The resulting hidden state is calculated by concatenating each hidden state output of the different heads and multiplying it with a contraction matrix that outputs a hidden state vector in its single-head dimensionality. Multi-head attention is needed to be able to better focus on multiple parts of the input sequence when encoding an input position, otherwise it might happen that most of the focus is set on the input position itself. [26] uses 8 heads. [4, 26]

Overall six layers of encoder and decoder components having individual weight matrices are stacked on top of each other to capture low-level as well as high-level features. The dimensionality of the hidden state is 512. Note that each position of the input sequence is encoded on an individual path through the transformer, which makes parallelization possible compared to LSTM based architectures. [4, 26]

The decoder uses almost the same architecture as the encoder, but contains an additional multi-head attention component that integrates all the hidden state encodings from the encoder. Therefore the final hidden state encodings resulting from the top most encoder are transformed into key and value attention vectors and given to the integrating multi-head attention component of every decoder. Once again this helps to better focus on the relevant input positions for the current output position based on hidden state information

extracted from the encoder. Note that the decoder works in a sequential manner again meaning that the input to the decoder are the already decoded sequence tokens of previous time steps, future input positions are masked away. The first multi-head attention component therefore capture inter-dependencies in the generated output sequence, the second one, the so called encoder-decoder attention, enriches the hidden state encoding with the information given from the encoder hidden states. Finally a linear layer mapping the hidden state vector to a logits vector with dimensionality of the given output dictionary and a *softmax()* function produce the output sequence using the greedy or beam search approach. The *EOS* token symbolizes the end of the output to be generated. [4, 26]

Input to the transformer are word embeddings also of fixed dimensionality of 512. Onto each word embedding a positional encoding following a specific pattern is added. These patterns are either learned or fixed and make sure that the distances of the input positions are projected onto the queue, key and value representations to be utilized during scaled dot-product attention. This also enables self-attention to be scaled to unseen lengths of sequences. Transformers usually contain an upper bound of sequence length to be processed, which is different to recurrent architectures that can process inputs of arbitrary length. Input sequences are usually padded up until the specified sequence length, the used *PAD* tokens are masked to be excluded from the self-attention components. [4, 26]

3 Approach

Figure 14 visualizes the steps of our approach, which are described in detail in the following chapters.

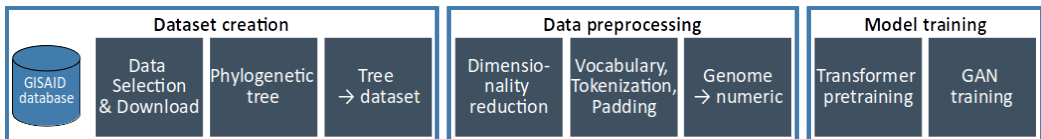


Figure 14: Approach pipeline [**own representation**]

3.1 Dataset Creation

As described in chapter 2.4 a dataset consisting of parent child pairs is needed to learn a ML model for mutation prediction. The steps to create this dataset are described in the following chapters.

3.1.1 Raw data selection from GISAID

In the first step of the dataset creation the raw genomes and their metadata are downloaded from GISAID. Therefore a selection of a suitable subpart of the data is necessary. The GISAID platform only allows the download of 5000 genomes at once. That is why we decided to focus our analysis on Germany (for one country the selection of < 5000 genomes is rather simple compared to the selection of data from multiple countries). By looking at the latest "Report on virus variants of SARS-CoV-2 in Germany" from the Robert Koch Institut (RKI) we decided for a time period. Figure 15 shows the distribution of different variants over time. Starting from calendar week 18 the Delta variant gradually displaces the previously widespread Alpha variant.

That is why we selected the data from 04.05.2021 (calendar week 18) until 15.07.2021 (calendar week 28). This results in a raw dataset of 35.818 genomes. Each genome consists of a fasta file containing the genomes sequence and a record in a tsv file with the metadata.

3.1.2 Generation of a phylogenetic tree

As described above we need parent child genome sequence pairs to be able to train a ML model. Therefore we need to evaluate the ancestral relationships between the genome sequences. As described in chapter 2.1.4 this is done in

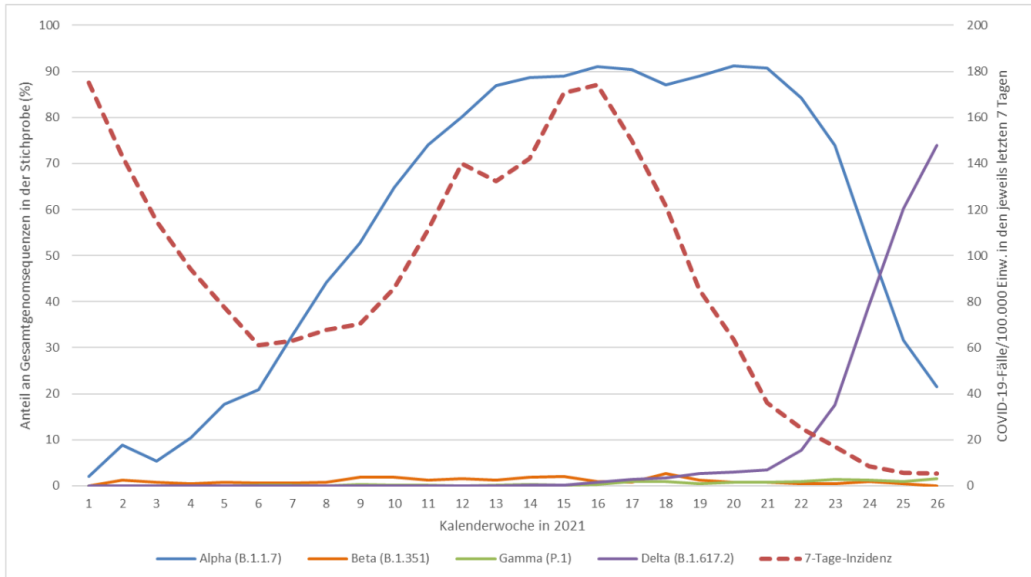


Figure 15: Variant distribution in Germany over time [3]

biology through phylogenetic trees. For the data preparation (e.g. alignment) and the calculation of the phylogenetic tree we use the nextstrain pipeline [11]. The pipeline was executed on a local computer with Intel Core i7-7500U (4*2.70GHz), NVIDIA GeForce 940MX and 16 GB RAM. Unfortunately after two days of calculation, the nextstrain pipeline exited with an out of memory exception. That is why the amount of data is decreased. This was done through iterative reduction of the dataset size. The largest locally computable dataset consists of 11.773 genome sequences.

The generated phylogenetic tree can be seen in figure 16.

3.1.3 Phylogenetic tree to dataset

Based on the calculated phylogenetic tree the final dataset is generated. Therefore the patristic distances between all leaf nodes are calculated. From the resulting patristic distance matrix for each leaf node the most related next leaf node can be evaluated. The two leaf nodes with the smallest distance are the closest relatives to each other and therefore represent a parent child pair. After the evaluation which leaf nodes are one parent child pair, the question which node is the parent and which node is the child is clarified. Therefore the two nodes are sorted by their date, which is stored in the metadata file. The older genome sequence is the parent and the younger genome sequence is the child.

For calculating the patristic distance matrix we first used Dendropy (version:

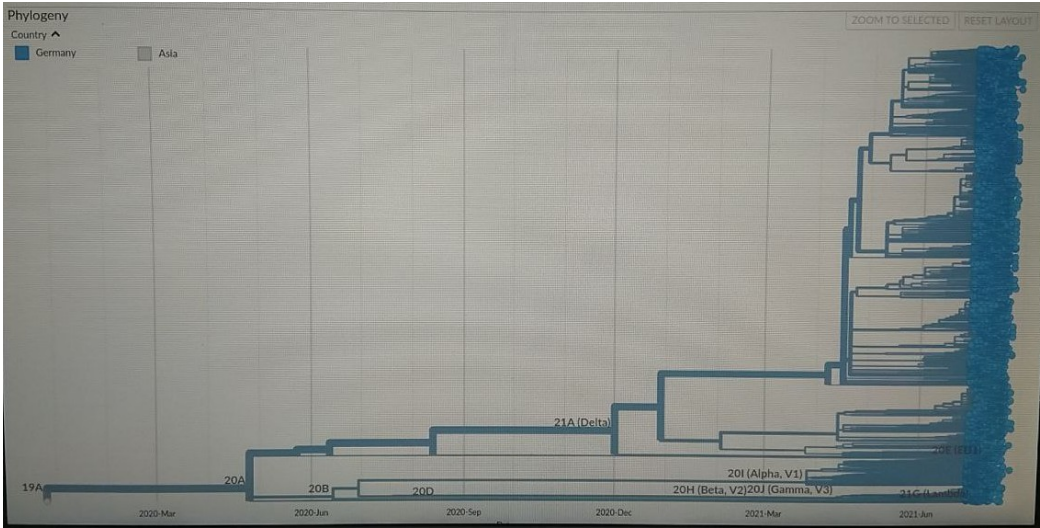


Figure 16: Phylogenetic tree of our dataset [own representation]

4.5.2) [9]. Again this leads to problems due to limited RAM on our local computer. As a consequence we switched to PhyloDM (version: 1.3.1) [18], which is a library for calculating patristic distances in a memory and time efficient manner.

Finally we received a dataset containing parent child genome sequences.

3.2 Data Preprocessing

3.2.1 Dimensionality reduction by selecting subpart of the genome

The full SARS-CoV-2 genome consists of about 30.000 nucleotides. Due to the limited computing capacity on our local computers we need to reduce the dimensionality of the dataset. This is done by selecting a subpart based on the gained data insights from chapter 4.1. In figure 18 parts of the genome with lots of mutations are visible. Based upon this we selected 99 nucleotides (33 amino acids) in the range between position 21.800 and 21.899.

3.2.2 Transform genome sequence to numeric model input

TODO

- Transform string to numeric for model input
- DNA Sequencing (Done during dataset creation, given from GISAID)
- DNA Sequence Tokenization for Amino Acid Dictionary

- DNA Sequence Padding

3.3 Model Architecture

TODO

3.4 Training Process

TODO

4 Experimental results

4.1 Dataset insights

This chapter presents the dataset insights of the complete dataset with full genome sequences and the dataset insights of the preprocessed dataset.

4.1.1 Complete dataset

The dataset consists of 9199 parent child pairs. The distribution how the parent and child sequences differ can be seen in figure 17. The majority of parent child pairs differ in less than 200 characters. The number of completely equal parent child pairs is 396.

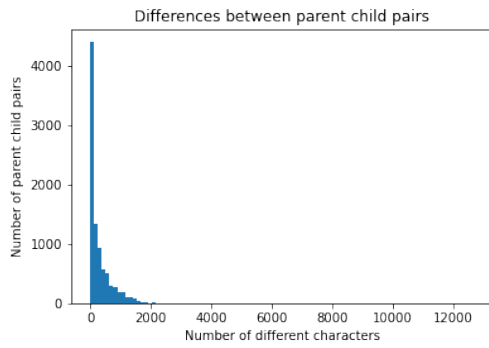


Figure 17: Distribution of the differences between parent and child sequences [own representation]

Figure 18 visualizes the distribution of mutations over the full SARS-CoV-2 genome.

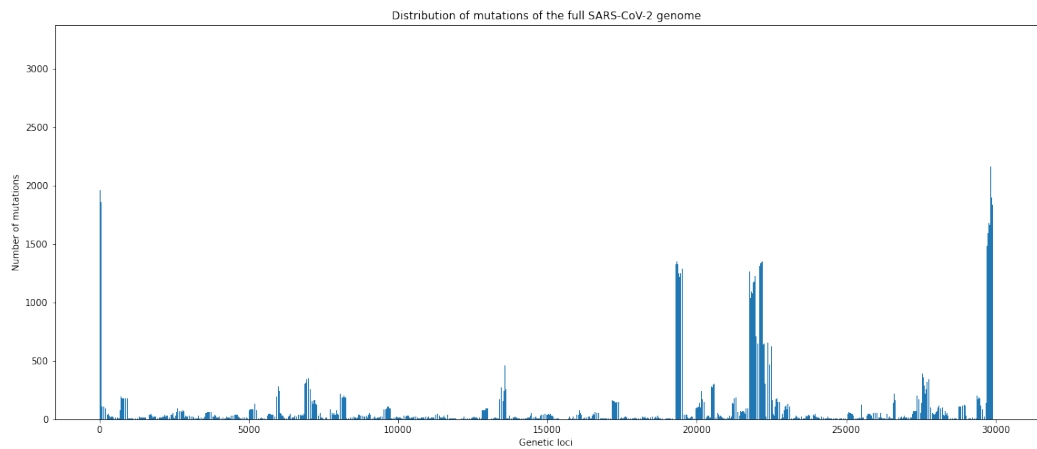


Figure 18: Mutated genetic loci [own representation]

4.1.2 Preprocessed dataset

The complete dataset is preprocessed as described in chapter 3.2.

The distribution how the parent and child amino acids differ in the selected subpart can be seen in figure 19. The majority (6946) of parent child pairs are equal.

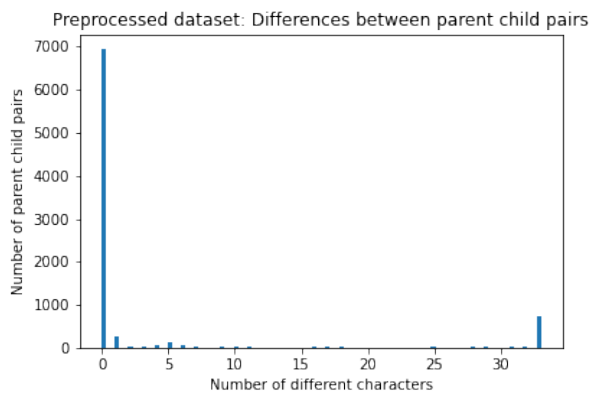


Figure 19: Preprocessed dataset: Distribution of the differences between parent and child sequences [own representation]

Figure 20 visualizes the differing amino acids and their positions of the selected subpart of the SARS-CoV-2 genome. They are distributed equally over the selected subpart.

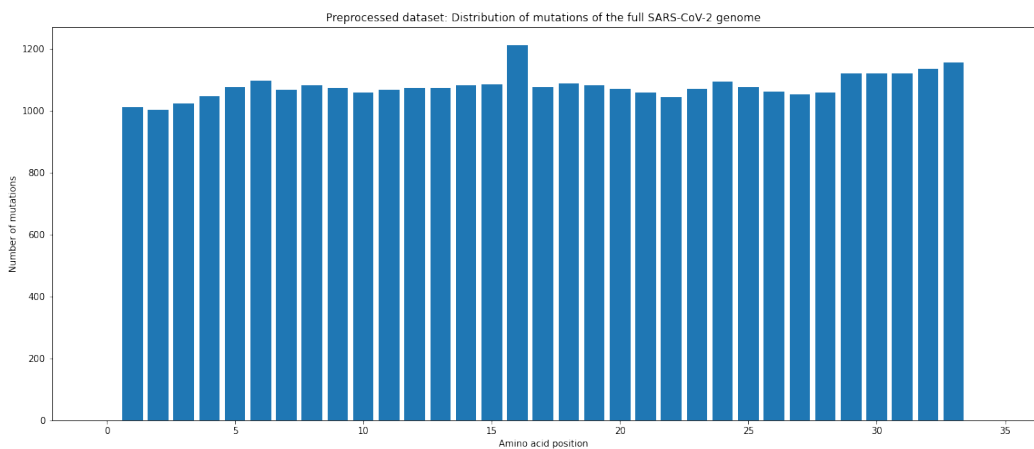


Figure 20: Preprocessed dataset: Differing amino acid positions [own representation]

4.2 Evaluation

For evaluating the model prediction we need to compare the generated sequence with the real sequence. For this text similarity task there are already metrics available in the NLP area. Berman et al. [6] have used the Bilingual evaluation understudy (BLEU) score. This metric comes from the area of machine translation and evaluates the similarity between machine generated text and human generated text. The more similar the two texts are, the better the BLEU score is. It is always in the range between zero and one. With one as perfect similarity and zero means no similarity.

4.2.1 Transformer pretraining

Figure 21 shows the loss plot for the Transformer pretraining. The model converges rather fast. The first meaningful results are already reached after 2 epochs. The reached BLEU score is 0.8559. From this point of view the results look very promising. That is why the details of these results are investigated. When comparing all predictions with the true values it is noticeable, that the model prediction is always the same for all 460 test instances. This could be explained by the fact, that the predicted sequence is correct in 318 of 460 cases. Therefore it can be assumed, that the model works, but the used dataset is not optimal.

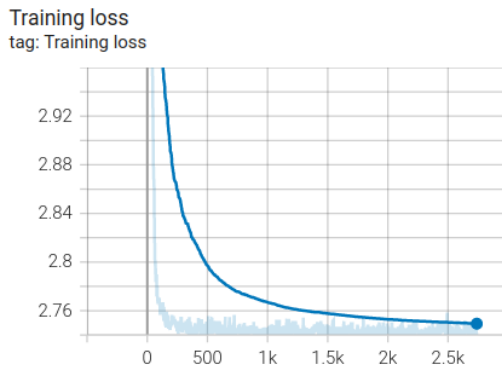


Figure 21: Transformer pretraining loss plot

4.2.2 GAN training

TODO

5 Conclusion

Coming to a conclusion, our main research question was whether a ML model can be trained to predict the next possible SARS-CoV-2 mutations.

The short answer to this is, that our trained ML model is definitely able to generate possible next sequences.

TODO: BLEU scores for pretraining and training TODO: is it really that convincing?

Furthermore, while answering this research question we created a new dataset based on raw data from GISAID. As part of this we developed a reusable pipeline for creating evolutionary datasets.

As an outlook, there are some improvements and further investigations we would like to evaluate in future work. As realized while generating data insights of the dataset, lots of our data instances are very similar to each other. To really model the worldwide development of SARS-CoV-2 mutation events, the raw dataset should be based on a subsampling of all available global data instances. Further improvements could be achieved by using a computing cluster for generating a larger dataset and for training the model on the full SARS-CoV-2 genome and not on a subset of the genome. Moreover in the model evaluation more domain specific metrics as proposed by Berman et al. [6] could be used in addition to the BLEU score.

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