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Strain Wars: Competitive interactions between SARS-CoV-2 strains are explained by Gibbs energy of antigen-receptor binding

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A B S T R A C T
Since the beginning of the COVID-19 pandemic, SARS-CoV-2 has mutated several times into new strains, with an increased infectivity. Infectivity of SARS-CoV-2 strains depends on binding affinity of the virus to its host cell receptor. In this paper, we quantified the binding affinity using Gibbs energy of binding and analyzed the competition between SARS-CoV-2 strains as an interference phenomenon. Gibbs energies of binding were calculated for several SARS-CoV-2 strains, including Hu-1 (wild type), B.1.1.7 (alpha), B.1.351 (beta), P.1 (Gamma), B.1.36 and B.1.617 (Delta). The least negative Gibbs energy of binding is that of Hu-1 strain, -37.97 kJ/mol. On the other hand, the most negative Gibbs energy of binding is that of the Delta strain, -49.50 kJ/mol. We used the more negative Gibbs energy of binding to explain the increased infectivity of newer SARS-CoV-2 strains compared to the wild type. Gibbs energies of binding was found to decrease chronologically, with appearance of new strains. The ratio of Gibbs energies of binding of mutated strains and wild type was used to define a susceptibility coefficient, which is an indicator of viral interference, where a virus can prevent or partially inhibit infection with another virus.

Introduction
In late 2019, a new infectious disease appeared in Wuhan, China, which later became known as COVID-19. Soon after, the disease agent was isolated, named SARS-CoV-2, and chemically (Popovic and Minceva, 2020; Popovic, 2022; Degueldre, 2021; Simšek et al., 2021) and thermodynamically characterized (Popovic & Minceva, 2020, 2021). The virus spread, causing a pandemic. The original strain was labelled Hu-1 (Islam et al., 2020). The pandemic has been active for 2 years, during which the virus has mutated several times. The strains have been labelled Hu-1 (wild type – Wuhan), B.1.1.7 (UK), B.1.351 (South Africa), P.1 (Brazil), B.1.36 (India, Canada and UK) and B.1.617 (India). The old strains are continuously suppressed by new strains. Obviously, interference has been taking place between various strains of SARS-CoV-2. Newer strains possess mutations giving them an advantage for survival. These changes are, from the chemical perspective, reflected in increased binding affinity of the mutant strains (Barton et al., 2021).

Infection is the result of interaction between a pathogen and its host organism, representing not only biological, but also a chemical and thermodynamic interaction (Kumar et al., 2021; Kruse et al., 2012; Popovic and Minceva 2020a, 2020b, 2021). Thus, due to this complexity, understanding infection demands a novel platform at the interface of virology, immunology, genetics, epidemiology, physical chemistry and biothermodynamics. If two pathogens simultaneously circulate in a population, the interaction becomes even more complex, as the pathogens compete not only with their host, but also with each other. When two viruses meet in a single host, the interaction is complex and threefold: each virus interacts with the host and the two viruses interact mutually (Popovic and Minceva, 2021). All three interactions are competitive, since both viruses and the host compete for a limited metabolic machinery and resources (i.e. amino-acids, nucleotides, energy sources etc.) (Popovic and Minceva 2020a, 2020b). Thus, the competition for resources represents the main mechanism of these interactions. The outcome of virus-virus-host interactions depends on two properties: susceptibility and permissiveness, both of which influence infectivity (Popovic and Minceva 2021). Susceptibility is the ability of a virus to enter the host cell (Duponchel and Fischer, 2019). On the other hand, once inside, permissiveness is the ability of a virus to multiply in the host cell (Duponchel and Fischer, 2019; Hou et al., 2017).

Infectivity is the ability of a pathogen to infect the host cell or organism. Infectivity varies between virus species. However, infectivity of SARS-CoV-2 differs between various strains of the same virus. Usually...
similar viruses do not show large differences in infectivity and perform coinfection (Popovic and Minceva 2021; Nickbakhsh et al., 2019). The infectivity of mutated strains of SARS-CoV-2 increases from the Wild type (Wuhan) to the new Delta strains (Thomas, 2021; Ramesh et al., 2021). What is the molecular basis for this trend? The evolutionary explanation states that this is the consequence of the tendency towards better adaptation to the host. This is the macroscopic explanation for the increased infectivity of the mutant strains. On the other hand, the microscopic explanation is based on changes on the receptor binding domain (RBD), making the antigen receptor binding more efficient. In microscopic explanation is based on changes on the receptor binding increased infectivity of the mutant strains. On the other hand, the better adaptation to the host. This is the macroscopic explanation for the phenomenon where a virus can prevent or partially inhibit infection with another virus within the same host (Schultz-Cherry, 2015; Wu et al., 2020; Dianzani, 1975).

A strain becomes dominant through competitive interaction with other strains, in the host population (Popovic and Minceva 2021). Obviously, an acquired mutation gives an advantage to one strain, enabling it to dominate. In extreme cases, domination leads to exclusion (Popovic and Minceva 2021). For example, the Iota strain was first discovered November 2020, and through interference came to represent 45% of new cases in February 2021 (Farinholt et al., 2021). Moreover, dual SARS-CoV-2 infection has been studied with two phylogenetically distant strains. The initially dominant strain belonged to GH clade and was suppressed by another strain from the GR clade, in only eight days (Samoilov et al., 2021). The initial ratio of the GH to GR strain was 70:30, while 8 days later it became 3:97, obviously as a result of interference (Samoilov et al., 2021). Thus, we have observed interference between various strains of SARS-CoV-2 (Korber et al., 2020). In that case a question is raised about the mechanism and driving force for this interference.

Gibbs energy represents the driving force of all chemical and physical processes in nature (von Stockar, 2013b; Demirel, 2014; Atkins and de Paula, 2011). Gibbs energy is important because it can be used to estimate the spontaneity and rate of a chemical process (von Stockar, 2013b; Demirel, 2014; Atkins and de Paula, 2011). One such process is antigen-receptor binding (Gale, 2020, 2019). Thus, Gibbs energy of binding is a very significant property of SARS-CoV-2 (Ngo et al., 2021).

The aim of this paper is to explore the mechanism and the driving force of interference between various strains of SARS-CoV-2 viruses. A thermodynamic approach will be used because the thermodynamic property, namely Gibbs energy represents the driving force for chemical reactions performed by viruses (von Stockar, 2013a; von Stockar, 2013b; Popovic and Minceva, 2021, 2020a, 2020b; Şimşek et al., 2021), enabling susceptibility and permissiveness. Virus-receptor binding and replication, transcription, translation and self-assembly are essentially chemical reactions driven by Gibbs energy. Faster virus multiplication leads to increase in the size of infective reservoir, causing the increase in

### Table 1

Gibbs energy of binding of SARS-CoV-2 strains. The values of the binding constant, $K_b$, and Gibbs energy of binding, $\Delta G$, values were calculated using $K_b$ values from the literature. Mutations led to increase in binding affinity and decrease in Gibbs energy of binding, implying greater spontaneity of binding of mutated strains. More negative Gibbs energy makes the process of antigen-receptor binding more favorable. The values have been calculated at 37°C (310.15 K).

| Date of isolation | PANGO lineage | WHO label | First outbreak | Mutations | $K_b$ (M⁻¹) | Reference | $\Delta G$ (kJ/mol) |
|-------------------|---------------|-----------|----------------|-----------|-------------|-----------|-------------------|
| Dec-19            | Hu-1          | Wild type | Wuhan          | Wild type | 2.13E-08    | Augusto et al., 2021 | 4.69E+07 | -45.55            |
| Dec-19            | Hu-1          | Wild type | Wuhan          | Wild type | 4.03E-07    | Ramanathan et al., 2021 | 2.48E+06 | -37.97            |
| 26-Jan-21         | B.1.1.7       | Alpha     | United Kingdom |           | 8.76E-08    | Ramanathan et al., 2021 | 1.14E+07 | -41.90            |
| May-20            | B.1.351       | Beta      | South Africa   |           | 2.04E-07    | Ramanathan et al., 2021 | 4.90E+06 | -39.72            |
| May-20            | B.1.351 and P.1 | Beta and Gamma | South Africa and Brazil | E484K | 1.97E-08 | Augusto et al., 2021 | 5.08E+07 | -45.75            |
| May-20            | B.1.351 and P.1 | Beta and Gamma | South Africa and Brazil | E484K | 9.90E-09 | Augusto et al., 2021 | 1.01E+08 | -47.53            |
| 10/1/2020         | B.1.617       | Delta     | India          |           | 4.60E-09    | Augusto et al., 2021 | 2.17E+08 | -49.50            |
| Dec-19            | Hu-1          | Wild type | Wuhan          | WT        | 6.26E-08    | Barton et al., 2021 | 1.60E+07 | -42.77            |
| 18-Dec-20         | B.1.1.7       | Alpha     | United Kingdom | N501Y (Alpha) | 5.3E-09     | Barton et al., 2021 | 1.82E+08 | -49.04            |
| 26-Jan-21         | B.1.351       | Beta      | South Africa   | K417N     | 3.49E-07    | Barton et al., 2021 | 2.87E+06 | -38.34            |
| 14-Jan-21         | B.1.351       | Beta      | South Africa   | K417N/E484K | 2.51E-07   | Barton et al., 2021 | 3.98E+06 | -39.19            |
| 14-Jan-21         | B.1.351       | Beta      | South Africa   | K417N/E484K/N501Y (Beta) | 1.74E-08 | Barton et al., 2021 | 5.75E+07 | -46.07            |
| Nov-20            | P.1           | Gamma     | Brazil         | K417T     | 2.62E-07    | Barton et al., 2021 | 4.42E+06 | -39.46            |
| Nov-20            | P.1           | Gamma     | Brazil         | K417T/E484K | 1.47E-07 | Barton et al., 2021 | 6.80E+06 | -40.57            |
| Nov-20            | P.1           | Gamma     | Brazil         | K417T/E484K/N501Y (Gamma) | 1.22E-08 | Barton et al., 2021 | 8.20E+07 | -46.99            |
infectivity. Influence of increased susceptibility has already been described in the literature (Ozono et al., 2021; Hasegawa et al., 2007).

Methods

A thermodynamic analysis is used to calculate Gibbs energies of binding of various strains of SARS-CoV-2. Gibbs energies of competing pairs of strains will be compared, to explain the phenomenon of interference of various pairs of virus strains.

Antigen-receptor binding represents a chemical process (Popovic and Minceva, 2021). The rate of this process is given by the phenomenological equation

$$ r_B = \frac{L}{T} \Delta G $$

(1)

Where $r_B$ is the rate of antigen-receptor binding, $L$ a constant (known as phenomenological coefficient), $T$ temperature and $\Delta G$ Gibbs energy of antigen-receptor binding (Popovic and Minceva, 2021; von Stockar, 2013a; Demirel, 2014; Hellingwerf et al., 1982; Westerhoff et al., 1982). The rate of binding is proportional to the absolute value of the Gibbs energy of binding. It is well documented that mutations lead to change in Gibbs energy of binding (Barton et al., 2021). The ability of coronaviruses to infect humans is invariably associated with their binding strengths to human receptor proteins (Zou et al., 2020). Mutation induces significant conformational transitions in the spike glycoprotein (Istifli et al., 2021). Natural selection promotes mutations that increase the spike ACE2 binding affinity (Istifli et al., 2021). Gibbs energy of binding can be calculated from dissociation constants, through the equation

$$ \Delta G = -RT \ln(K_B) $$

(2)

Where $R$ is the universal gas constant and $T$ temperature (Popovic and Minceva, 2021; Du et al., 2016). $K_B$ represents the binding constant, which is the reciprocal of the dissociation constant $K_D$ (Du et al., 2016).

$$ K_B = \frac{1}{K_D} $$

(3)

The dissociation constant is the equilibrium constant of the dissociation reaction of the antigen-receptor complex into the receptor and antigen

$$ RA + R = A $$

(4)

where $R$ represents the host cell receptor (ACE2), $A$ the virus antigen (spike protein) and $RA$ the receptor antigen complex (Du et al., 2016). Thus, $K_B$ is defined as the ratio of concentrations of the free receptor [$R$] and antigen [$A$] to the receptor antigen complex [$RA$] (Du et al., 2016)

$$ K_D = \frac{[R][A]}{[RA]} $$

(5a)

However, $K_D$ can also be defined through kinetic parameters of the reaction: the association, $k_on$ and dissociation, $k_off$ rate constants.

$$ K_D = \frac{k_{on}}{k_{off}} $$

(5b)

Dissociation constants of various SARS-CoV-2 strains have been reported by (Augusto et al., 2021; Barton et al., 2021; Ramanathan et al., 2021) and are given in Table 1. The reported data are for binding between a single antigen (SGP trimer) and a single receptor (ACE2).

The binding constant, $K_B$, was calculated using Eq. (3), from $K_D$. The dissociation constant refers to reaction (4), between a single antigen (SGP trimer) and a single receptor (ACE2). Thus, the calculated $K_B$ is for a single antigen-receptor interaction. This might not be the case if there are multiple SGP-trimer/ACE2 interactions, during virus binding (Gale, 2021). In that case, $K_B$ for the entire virus might be much greater than that for a single SGP trimer/ACE2 binding (Gale, 2021). Moreover, $K_B$ will be reduced if entropy decreases during whole-virus binding (Gale, 2021). However, human coronaviruses share a very similar size and structure (Neuman and Buchmeier, 2016). Thus, an assumption can be made that the number of SGP/ACE2 interactions is the same for all SARS-CoV-2 strains and that the entropy change on virus binding is the same for each strain (Gale, 2021). In that case, $K_B$ for the SGP/ACE2 interaction is proportional to, or is at least an indication of, the relative magnitudes of $K_B$ for the whole virus (Gale, 2021). This is because the $\Delta G$ values are additive for each SGP/ACE2 interaction (Gale, 2021), meaning that multiple SGP/ACE2 interactions will not change the conclusions of the paper.

Finally, we will introduce a new property – susceptibility coefficient $\tilde{S}$, which is defined as the ratio of rates of binding of two viruses or in this case a mutant strain and wild type virus.

$$ S = \frac{r(\text{mutant})}{r(\text{wild type})} $$

(6)

Combining this equation with the phenomenological Eq. (1) gives

$$ S = \frac{\Delta G(\text{mutant})}{\Delta G(\text{wild type})} $$

(7)

The $I$ coefficient and $T$ are equal for both strains, since both attack the same host at the same physiological temperature. The phenomenological coefficient is proportional to the equilibrium forward and backward half-reaction rate (Demirel, 2014). These in turn depend on the forward and backward the association, $k_{on}$ and dissociation, $k_{off}$ rate constants (Du et al., 2016). Since $k_{on}$ and $k_{off}$ vary between different virus strains (Barton et al., 2021), it might be possible that $I$ differs between virus strains. This will be a subject of our future research.

Results

Based on dissociation constants, Gibbs energies of binding were calculated for various SARS-CoV-2 strains. The results are presented in Table 1. The analyzed SARS-CoV-2 strains include Hu-1 (wild type), B.1.1.7 (alpha), B.1.351 (beta), P.1 (Gamma), B.1.36 and B.1.617 (Delta). The original Hu-1 strain has been designated as wild type and was isolated in Wuhan, China in December 2019. The strain B.1.1.7 has been designated Alpha and was first isolated in the United Kingdom in late January 2021. The strain B.1.351 was named Beta, isolated in South Africa in May 2020. The strain P.1. was labeled Gamma and was first isolated Brazil in November 2020. The strain B.1.36 has been initially reported from India, Canada, and UK (Basheer and Zahoor, 2021). Finally, the strain B.1.617 has been labeled Delta and was first isolated in India, in October 2020.

The analyzed strains exhibit a variation in binding constants and hence Gibbs energies of binding according to Eq. (2). The B.1.1.7 (Alpha) has the most negative Gibbs energy of binding, of $-50.06$ kJ/mol. On the other hand, the least negative Gibbs energy of binding is that of Hu-1 strain, $-37.97$ kJ/mol. Moreover, Gibbs energies of binding decrease chronologically, with appearance of new strains. Similarly, the binding constants span the range between $2.48 \times 10^9$ M and $2.70 \times 10^9$ M. All the data in Table 1 have been calculated at the physiological temperature of $37^\circ C$ ($310.15$ K).

Discussion

We hypothesize that mutations that appeared during time have led to increase in binding constant and more negative Gibbs energy of binding. According to the phenomenological Eq. (1), this leads to a greater binding rate, which in turn leads to more rapid cell entry and multiplication of one of the strains. Finally, this results in a greater infectious reservoir and greater infectivity.

Virus-virus interactions influence the epidemiology of respiratory infections (Dee et al., 2021; Popovic and Minceva 2021). After the initial
development of the pandemic caused by the Hu-1 wild type strain, mutations have developed with time, in various countries. Chronologically, pairs of strains compete for resources, through indirect interaction of strain pairs (Popovic and Minceva, 2021). Thus, the discussion will be represented as an arena, where the strains compete. In the beginning, the Hu-1 strain spread across the planet and had no competition, since no other strains existed. With the appearance of the Alpha and other strains, except for the direct interaction with the host, an interaction between two strains of SARS-CoV-2 occurred.

Wild type vs Alpha (Hu-1 vs B.1.1.7)

The Hu-1 strain is characterized by a Gibbs energy of binding of $-37.97 \text{ kJ/mol}$. Simultaneously, the mutated B.1.1.7 strain is characterized by a Gibbs energy of binding of $-41.90 \text{ kJ/mol}$. Several authors reported various values of dissociation constants for the same strain. Thus, several values can be found in the table. Having in mind that Gibbs energy of the B.1.1.7 strain is more negative, one can conclude that the binding rate for this strain will be greater. Thus, during competition, the strain B.1.1.7 will enter the host cell faster and thereby gain an advantage while hijacking the host cell metabolism. This advantage will lead to suppression of the Hu-1 strain.

Wild type vs Beta (Hu-1 vs B.1.351)

In South Africa, simultaneously in circulation, appeared the Wild type and Beta strains. From epidemiological studies, it is known that there had been interference and the Wild type strain was suppressed...
Thus, we can expect that Gibbs energy of the Beta strain will be more negative than that of the Wild type. Indeed, Gibbs energy of binding of the Wild type is \(-37.97\) kJ/mol, while that of the Beta strain is \(-39.72\) kJ/mol. Like in the previous case, due to more negative Gibbs energy of binding, the Beta strain has an advantage to enter the host cell faster and hence multiplies faster and increases the infectious reservoir, making Beta strain more infective. 

Wild type vs Gamma (Hu-1 vs P.1) 

In Brazil, simultaneously in circulation, appeared the Wild type and Gamma strains. The Wild type has a Gibbs energy of binding is \(-37.97\) kJ/mol. The Gamma strain has a Gibbs energy of binding of \(-39.46\) kJ/mol. Since the Gamma strain has a more negative Gibbs energy of binding and greater affinity for the receptor, it enters the host cell and multiplies faster, leading to interference and suppression of the Hu-1 strain.

Wild type vs Alpha vs Delta (Hu-1 vs B.1.1.7 vs B.1.617) 

In 2021, in Europe, three strains met simultaneously. Through the same competition mechanism, the strain with the most negative Gibbs energy of binding will have the greatest rate of binding, enter the cell the fastest and multiply the most rapidly. This leads to the interference phenomenon, resulting in suppression of the two other strains. Indeed, the Wild type has a Gibbs energy of \(-37.97\) kJ/mol, the Alpha strain has a Gibbs energy of binding of \(-41.90\) kJ/mol, while the for the Delta strain it is \(-49.50\) kJ/mol. The Delta strain has the most negative Gibbs energy of binding. Thus, it was able to suppress the other two strains. Notice that every wave of the pandemic has been followed by appearance of a new strain. Every new strain had a greater infectivity and hence the number of infected grew with each wave of the pandemic. Newly emerging variants of SARS-CoV-2 have contributed to successive waves of COVID-19. The new variants increased disease severity and viral transmissibility, thus increasing the morbidity and mortality of COVID-19 (Thomas, 2021; Ramesh et al., 2021). The increase in wave size is the consequence of faster entry of the virus into its host cells and multiplication inside the cell.

The anti-epidemic measures of wearing protective masks, distancing, isolation of the diseased and their contacts, as well as vaccination, are making a selective pressure on the virus. All these measures attempt to decrease the infectious reservoir and the possibility of its transmission. The virus reacts to this selective pressure by evolving towards greater infectivity and transmission rate. This is reflected in increase in binding affinity and decrease in Gibbs energy of binding. The strains are not fighting each other, but each strain tends to develop a more efficient mechanism for its survival. Natural selection can act upon rare but favorable mutations (Korber et al., 2020). This unfortunately means that the fight against SARS-CoV-2 will have to continue with development of new vaccines and medicines, adapted to new strains.

Conclusion 

The basic mechanism of the interaction between various strains of SARS-CoV-2 is competition. The competition is reflected at the susceptibility and permissiveness levels. This paper analyzes only susceptibility, since only data on binding constants have been available in the literature. It would be possible to determine the permissiveness for various strains, if their elemental composition were known. This is unfortunately currently not the case. Since the counter-epidemic measures are making a selective pressure on the virus, the virus evolves towards an increased binding affinity and more negative Gibbs energy. This has been shown by the data given in Table 1. Gibbs energies of binding decrease chronologically, with appearance of new strains. The Delta strain has the most negative Gibbs energy of binding. SARS-CoV-2 is also expected to evolve towards a changed permissiveness, also through mutations on other parts of the viral nucleic acid. The test of this hypothesis can be made in the future, once the elemental composition of the SARS-CoV-2 strains is known.

Conflict of interest statement 

The authors declare no conflict of interest.

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