

Research Article

A Population-Based Model for Rationing COVID-19 Vaccine

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Background

As COVID-19 vaccines develop, methods for identifying vulnerability within groups to prioritized vaccination remain un-established. This paper presents a novel approach based on population-based analysis of viral pneumonia vulnerability, as an example.

Methods

The analysis employed an anonymous, 16-year, population dataset (n = 768,460) consisting of International Classification of Diseases (ICD-9) diagnoses, demographics, and dates identifying those with viral pneumonia and permitting linkage of these individuals to all their associated diagnoses for calculation of odds ratios and proportions of disorders before and after the index viral pneumonia diagnosis.

Results

Females and males had results of differing magnitude. For those with viral pneumonia, the mean number of diagnoses was greater in both the subsample and whole sample, with associated diagnoses arising about 4 years on average before the viral pneumonia index diagnosis. Within the subsample, compared to those without, the temporal analysis revealed distinct over-representation for those with viral pneumonia at visit one and over the first fifty visits. Further, those with viral pneumonia had diagnoses not represented in the group without viral pneumonia.

Conclusions

The population-based analysis of temporal hyper-morbidity may be a viable and economical approach to identifying viral pneumonia vulnerability. The approach presented in this paper may provide an economical means of identifying vulnerability to COVID-19 in regions where comparable data are available for analysis. Rational approaches may optimize vaccination and help to limit the spread of the disease and to some extent alleviate the health service burden.

Introduction

With the advent of COVID-19 vaccines and given a relatively limited initial supply comes the requirement for decisions regarding who first receives vaccination to best curb the spread of the pandemic. Vulnerable groups are identified based on infection rates and mortality, such as healthcare workers and the elderly. Yet, it is within groups across populations that there are presently no clearly defined criteria to serve as a basis on which to optimally ration vaccine. This paper presents for consideration one possible set of criteria not yet described in the literature.

Most models predicting viral pneumonia tend to focus on surveillance data.^{[1][2][3][4][5]} Others tend to focus on aspects of the virus species^{[6][7]} or variants within species.^[8] This paper presents a novel model based on the analysis of diagnosis data readily available in many health care catchments. The model derives from the population-based examination of temporal hyper-morbidity. The approach to morbidity analysis is an emerging field of investigation.^[9] For example, the World Psychiatry Association has recently established a comorbidity section based, in part, on work describing the population-based, temporal, hyper-morbidity of psychiatric disorder in relationship to other disorders, such as cancer and ulcerative colitis.^{[10][11]} This analysis employed a similar standardized, population-based approach to model development. The present model describes viral pneumonia-associated morbidity and provides an example of the temporal hyper-morbidity for all observed disorders in a subsample of those aged less than one year of age who subsequently did or did not develop viral pneumonia.

Method

In Alberta Canada, all physicians must bill the provincial government for reimbursement of each patient visit and record at minimum a patient identifier, date of birth, a diagnosis, and visit date. This study employed an anonymous 16-year population dataset (April 1993–November 2010) consisting of International Classification of Disease (ICD version 9) diagnoses, age, sex, and visit date for all health-seeking individuals in the Calgary Health Zone, Alberta Canada (ethics IDREB15–1057). The sample (Table 1) was stratified on the basis of sex, age (overall and those less than one year of age), and grouped on the presence (+) or absence (–) of any viral pneumonia (VP). Further, the encrypted unique identifier permitted linking as a group all diagnoses of those with viral pneumonia (VP+).

Data Analysis

To examine the overall relationship of the presence or absence of viral pneumonia (VP+ or VP-) and all other main classes of ICD-9 diagnoses in the study population, odds ratios and upper and lower 95% confidence intervals (CI) were calculated separately for females and males (Tables 2 a and b).

To provide an example for representation, a subsample of those under the age of one year was isolated (Table 1). This subsample was ordered by date of each visit and diagnosis with the frequency of each diagnosis tallied by order of visit. Figures 1 a and b show visit #1 for each sex, respectively, the ratio of the total frequency for each diagnosis divided by the sample sizes (Table 1) for those with (numerator: VP+) and without (denominator: VP-) viral pneumonia (VP). The horizontal y-line at the value one demarcating the proportions with values greater than one that indicate a greater proportional frequency of all ICD diagnoses in the numerator (VP+). Figures 2 a and b show the same calculation in three dimensions for the first 50 visits of those under the age of one year. The x-z lines at values one similarly demarcate equal ratios of proportions of diagnoses between the groups with greater frequency proportions within the linked VP+ diagnoses. Figures 2 a and b truncate at the value 4 for ease of peak and trough comparison, noting that the plateaus signify diagnoses where the [VP+/VP-] ratio is greater than the value 4, with the full distributions described in the text. Note that V codes were sequential ordered in the 1200 range with the value 12 replacing the letter V. Further, visits without diagnoses representing pathology, laboratory, or procedures were coded with the value 1300 for graphical representation, with the total noted in the subscript (*) of Table 1.

Results

Table 1 describes the for males and females of all ages and for those less than one year of age the counts of unique VP+ and VP- individuals and linked diagnosis frequencies, as well as means with standard deviations (SD).

Groups		Female	Male
All Ages			
VP-	UID	304505	267498
	Diagnoses*	36510566	20255378
	mean (SD)	120 (125)	76 (99)
VP+	UID	111821	84636
	Diagnoses*	25082428	13998139
	mean (SD)	224 (196)	165 (181)
< 1 Year			
VP-	UID	12775	20326
	Diagnoses*	139689	249063
	mean (SD)	11 (9)	13 (10)
VP+	UID	5125	8867
	Diagnoses*	69178	138234
	mean (SD)	14 (12)	16 (19)

Table 1. Group descriptions.

** Included 11,629,494 unspecified non-diagnosis counts for Pathology/Laboratory/Procedures*

Table 2 a and b show by sex the cell sizes (a, b, c, d) and odds ratio (OR), upper and lower 95% confidence intervals for each ICD main class in descending order. In each ICD-9 main class, the lower 95% confidence interval is greater than the value one, indicating over-representation of VP+-linked disorders within ICD classes for each sex. The lower limit of the 95% confidence intervals was greater in males than females in the following main ICD classes: neoplasms, blood/blood organs, complications of pregnancy, congenital anomalies, perinatal, and HIV. The upper limit of the 95% confidence intervals was in males were less than that of females in the following main ICD classes: endocrine, etc., mental

disorders, nervous system/sense organs, digestive system, skin, and subcutaneous tissue, musculoskeletal system connective tissue, ill-defined conditions, injury and poisoning, V codes, mental disorder combined with relevant V codes, and other respiratory diseases. The overall order of main ICD classes was similar for females and males, with the greatest magnitude being in males for diseases and disorders of blood/blood organs and with males having had greater complications of birth and pregnancy (e.g., fetal distress).

ICD Main Class	a	b	c	d	OR*	Lower 95%CI	Upper 95%CI
Ill Defined Conditions	33209	271296	2175	109646	6.17	5.91	6.45
Respiratory System	58683	245822	4600	107221	5.56	5.4	5.74
Other Respiratory Diseases	65105	239400	6778	105043	4.21	4.11	4.33
Nervous System/Sense Organs	90345	214160	12098	99723	3.48	3.41	3.55
Injury And Poisoning	78576	225929	10508	101313	3.35	3.28	3.43
V Codes	31778	272727	3794	108027	3.32	3.21	3.43
Skin And Subcutaneous Tissue	93041	211464	15042	96779	2.83	2.78	2.88
Musculoskeletal System Connective Tissue	93513	210992	15388	96433	2.78	2.73	2.83
Digestive System	163436	141069	34957	76864	2.55	2.51	2.58
Infectious/Parasitic	131878	172627	26536	85285	2.46	2.42	2.49
Mental Disorder with associated V Codes	118534	185971	23488	88333	2.4	2.36	2.44
Mental Disorders	124260	180245	25358	86463	2.35	2.31	2.39
Circulatory System	189254	115251	48397	63424	2.15	2.12	2.18
Genitourinary System	75375	229130	14783	97038	2.16	2.12	2.2
Endocrine, Nutritional, Metabolic Immune System	192767	111738	50810	61011	2.07	2.04	2.1
Blood/Blood Organs	258772	45733	82741	29080	1.99	1.96	2.02
HIV	198777	105728	58261	53560	1.73	1.7	1.75
Neoplasms	193759	110746	57012	54809	1.68	1.66	1.71
Congenital Anomalies	287414	17091	102593	9228	1.51	1.47	1.55
Perinatal	283998	20507	102294	9527	1.29	1.26	1.32
Complications of Pregnancy	224084	80421	80766	31055	1.07	1.06	1.09

Table 2a. Female Odds Ratios of +/- VP by ICD Main Classes.

$$*Odds\ Ratio = [(a*d)/(c*b)]$$

ICD Main Class	a	b	c	d	Odds Ratio	Odds Ratio	Lower 95%CI
Respiratory System	68105	199393	5135	79501	5.29	5.13	5.45
Ill Defined Conditions	45616	221882	3479	81157	4.8	4.63	4.97
Other Respiratory Diseases	75742	191756	7657	76979	3.97	3.87	4.07
Nervous System/Sense Organs	96971	170527	13395	71241	3.02	2.96	3.09
V Codes	65758	201740	8531	76105	2.91	2.84	2.98
Injury and Poisoning	61682	205816	8410	76226	2.72	2.65	2.78
Blood/Blood Organs	248851	18647	70507	14129	2.67	2.61	2.74
Skin and Subcutaneous Tissue	103086	164412	16386	68250	2.61	2.56	2.66
Infectious/Parasitic	135413	132085	24452	60184	2.52	2.48	2.57
Digestive System	155592	111906	30573	54063	2.46	2.42	2.5
Musculoskeletal System Connective Tissue	97838	169660	17146	67490	2.27	2.23	2.31
Mental Disorder with Associated V codes	138296	129202	27905	56731	2.18	2.14	2.21
Mental Disorders	143604	123894	29660	54976	2.15	2.11	2.18
Genitourinary System	186020	81478	43818	40818	2.13	2.09	2.16
Circulatory System	183415	84083	43226	41410	2.09	2.06	2.12
Endocrine, Nutritional, Metabolic Immune System	188328	79170	46009	38627	2	1.97	2.03
Complications of Pregnancy	264140	3358	82511	2125	2.03	1.92	2.14
HIV	187453	80045	47525	37111	1.83	1.8	1.86
Neoplasms	202394	65104	53390	31246	1.82	1.79	1.85
Perinatal	259121	8377	79994	4642	1.79	1.73	1.86
Congenital Anomalies	255067	12431	78360	6276	1.64	1.59	1.7

Table 2b. Male Odds Ratios of +/- VP by ICD Main Classes

Figure 1 shows for females (upper) and males (lower) the visit #1 [(VP+/VP-] ratios of sample proportions for those with and without VP for each diagnosis. The ratios of diagnoses with over-representation within the linked diagnoses of the VP+ group is given when the dropline ends above the horizontal y line (value one).

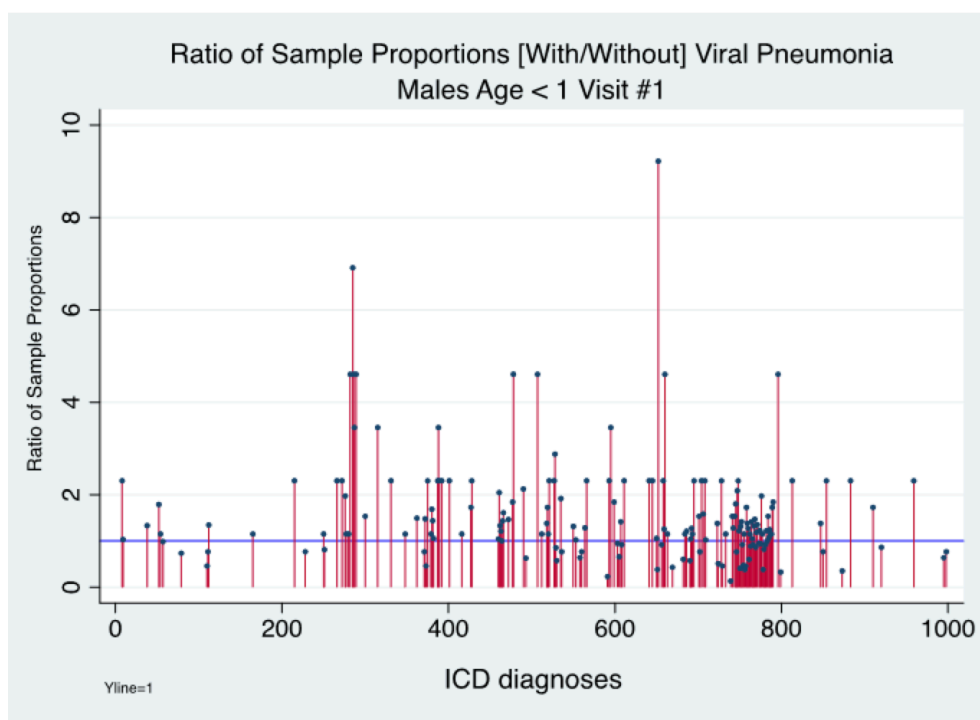
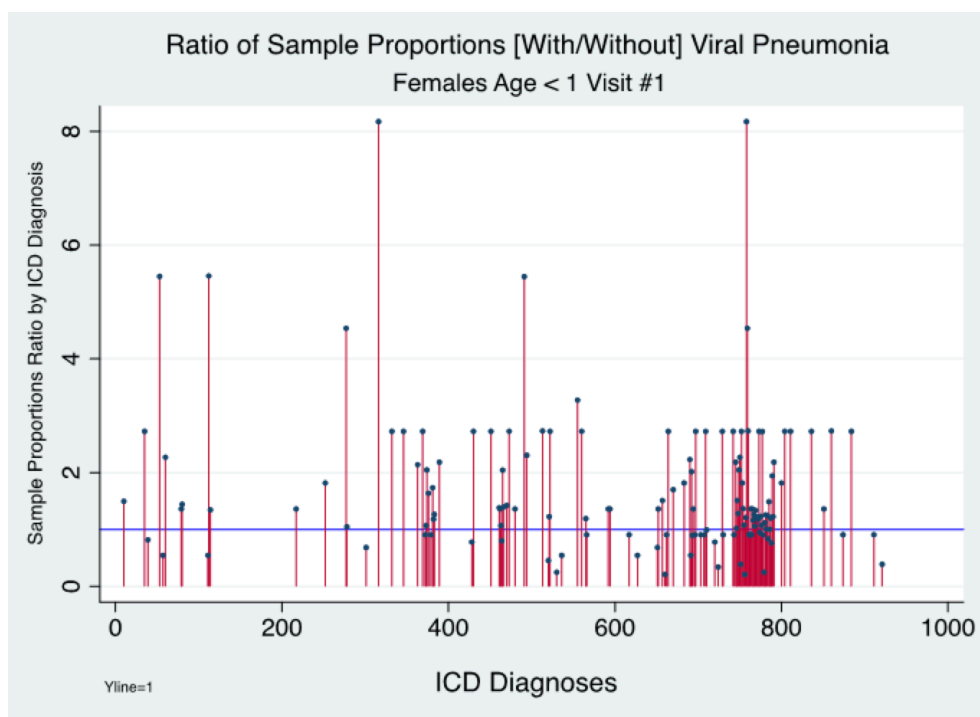


Figure 1. Sample proportions of female (upper) and male (lower) visit #1 diagnoses before index viral pneumonia diagnosis.

There were ICD diagnoses for which the VP+ group was assigned a diagnosis on visit #1 but for which there was no corresponding VP- diagnosis on which to base a comparison. Ratios were not calculated for

ICD-9 diagnoses not represented in one or other of the VP+ or VP- groups. The female VP- group for visit #1 contained 49 diagnoses not assigned in the VP+ group (not listed). Similarly, the female VP+ group contained 10 diagnoses not assigned in the VP- group: The ICD codes unique to the VP+ group were as follow: 345, 385, 426, 629, 705, 772, 793, 882, 953, and V code 5.

The male VP- group for visit #1 contained 130 diagnoses not assigned in the VP+ group (not listed). The male VP+ group contained 36 diagnoses not assigned in the VP- group: The ICD codes unique to the VP+ group were as follow: 1, 41, 190, 210, 230, 239, 259, 308, 323, 335, 345, 365, 377, 385, 426, 432, 455, 514, 523, 537, 579, 629, 653, 668, 698, 705, 772, 793, 820, 882, 945, 953, 999, V code 3, V code 5, and V code 50.

Figure 2 a and b show for each sex the three-dimensional variations in the ratios over time for all [VP+/VP-] linked diagnosis frequencies represented in temporal order by all ICD diagnoses for the first 50 visits for those under the age of one year. The Figure 2 graphics extends visit #1 ratios shown in Figure 1 across the first 50 visits for the less than one year age group (e.g., no visits for those age 1 or older are included). As noted in the methods section, the upper limit of the ratio for each diagnosis is truncated at the value four for ease of comparison. The visits were truncated at 50 for ease of visualizing variations across the range of represented ICD diagnoses.

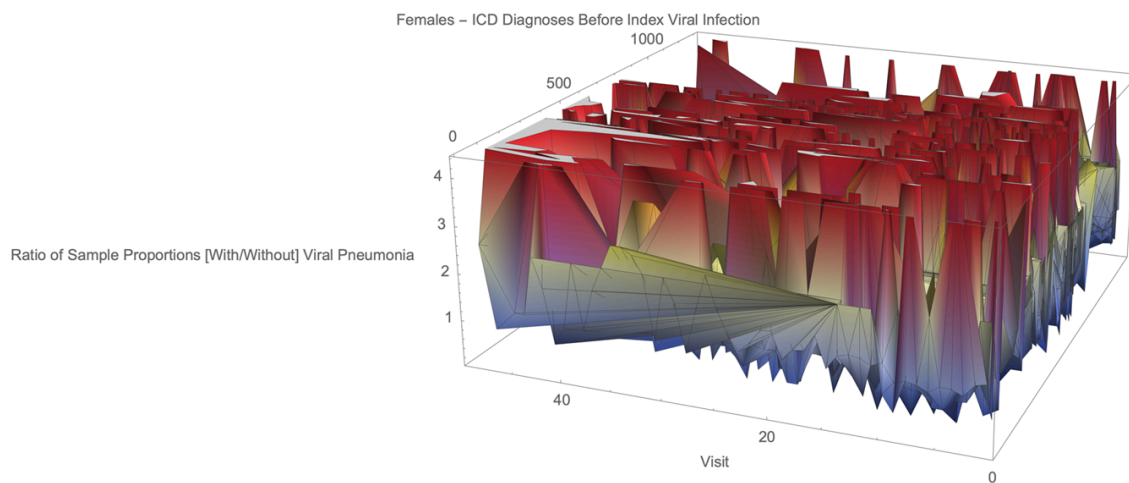


Figure 2a. Sample proportion ratios of female diagnoses for visits #1 to #50 in sequence before index viral pneumonia diagnosis.

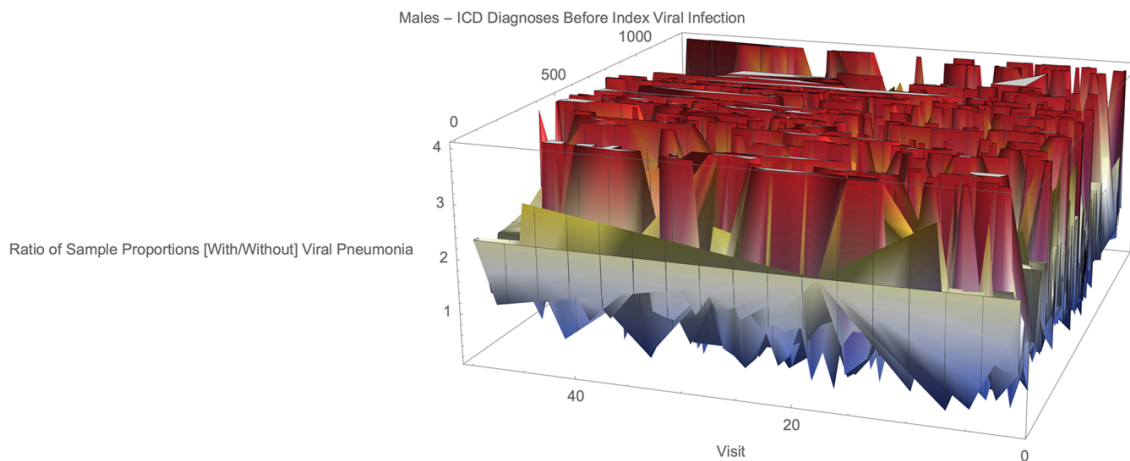


Figure 2b. Sample proportion ratios of male diagnoses for visits #1 to #50 in sequence before index viral pneumonia diagnosis.

The means, standard deviations (SD), and ranges of the full distributions of diagnosis ratios are described as follows. In the female group with age less than one year, there were 175 ICD diagnoses on average 3.97 (SD 3.65) years before VP+ where the ratio was greater than the value one with mean 1.43 (SD 1.31) and an upper range limit of 16, with 70 ICD diagnoses where the ratio was greater than the value 1.25 with mean 1.91 (SD 2.0). For females of all ages, ICD disorders preceded VP+ on average by 4.26 (SD 3.60) years.

In the male group with age less than one year, there were 310 ICD diagnoses on average 3.76 (SD 3.56) years before VP+ where the ratio was greater than the value one with mean 1.4 (SD 0.75) and an upper range limit of 9, with 166 ICD diagnoses where the ratio was greater than the value 1.25 with mean 1.74 (SD 0.93). For all ages, ICD disorders preceded VP+ for males on average by 4.19 (SD 3.58) years.

To summarize, not surprisingly, compared to those without, those with viral pneumonia have greater morbidity on average at all ages and when only those under the age of one year are considered (Table 1). There was an overall relationship between viral pneumonia (compared to those without) and the main ICD classes in the study population based on the odds ratio calculations shown in tables 2a and 2b. The odds ratio analysis was based on counts of individuals and did not provide information about the frequency (intensity) of the unique diagnoses within individuals making up each group. For both age groups, the ICD 9 diagnoses arising before the index viral pneumonia diagnosis did so by about 4 years on average. As such, the profile of diagnoses in the less and one year of age group would not on average have developed viral pneumonia until age four to five.

Examining the sequential diagnoses of those less than one year of age provided evidence of distinct profiles of diagnosis intensity comparing those with and without viral pneumonia when diagnoses were represented in both groups. When diagnoses were not represented in both groups, the analysis identified additional diagnoses unique to those with and without viral pneumonia.

Discussion

The population-based comparative analysis demonstrated an overall relationship of viral pneumonia across all ICD classes of disease. Analysis of the temporal order revealed an age-specific range of ICD diagnoses distinguishing those who subsequently developed viral pneumonia. The temporal analysis focused on ICD diagnoses preceding viral pneumonia, identifying the ICD diagnosis profiles of those vulnerable to viral pneumonia. This vulnerability, exemplified in a subsample less than one year of age on visit one (diagnosis) and over the first 50 visits (diagnoses), likely persists across the full range of age groups in the dataset given the greater average frequency of diagnoses in the VP+ group and the consistent period of time that associated ICD disorders precede viral pneumonia in the whole sample.

Where similar data exist at local, regional, and national levels, applying this population-based comparative analysis of temporal hyper-morbidity approach provides a standardized means of identifying those who have to date become infected with COVID-19. Identifying this vulnerability in those not yet infected may assist in prioritizing the most vulnerable for vaccination; an action that may more rapidly alleviate the social burden brought on by the COVID-19 pandemic.

There were several limitations in the present study. Diagnostic precision is a limitation, as not all physicians have the same level of practice competence with respect to the diagnostic formulation. For brevity, the subsample analysis focused on the ICD diagnoses preceding viral pneumonia. Focusing on associated hyper-morbidity following the index viral pneumonia may assist health systems to plan for increased demand of specific service types typical of this viral pneumonia. Lastly, while the sequential order of the first fifty visits was described, this order did not take into account the conditional orders of diagnoses within sets of visits (e.g., given first diagnosis X and second diagnosis Y, etc.). Compared to past analyses, the present analysis somewhat advances the method in terms of representing sequential diagnoses related to visits in time. However, taking into account the varying conditional orders temporal order of diagnoses leading to viral pneumonia within clusters of individuals may reveal more specific profiles of vulnerability with much greater precision.

In conclusion, there are many pathways of vulnerability to human viral pneumonia in addition to simple transmission from vector. Human genetic vulnerability^[12], while presently rare, is one pathway. Discovering and understanding the complex relationships within and between fields of genetics and metabolomics research hold great potential to prevent and cure, as these relate to viral infection in general and COVID-19 infection specifically.^{[13][14]} However, much integration of existing knowledge into practical application from across these fields of study into health care remains to be accomplished, even in respect to the 1918 influenza pandemic.^[15] The present work presents COVID-19-relevant information structured in a rapidly reproducible and universally applicable form to potentially help to curb the COVID-19 pandemic.

Statements and Declarations

Contributions: David Cawthorpe conceptualized, developed, executed the study and wrote this paper.

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Declarations

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