Race and Sex Differences in Vital Signs Associated with COVID-19 and Flu Diagnoses in Mississippi

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Abstract

Early detection of viral infections, such as COVID-19 and flu, have potential to reduce risk of morbidity, mortality, and disease transmission through earlier intervention strategies. For example, detecting changes in vital signs have the potential to more rapidly diagnose respiratory virus diseases. The objective of this study was to utilize the University of Mississippi Medical Center’s extensive clinical database (EPIC) to investigate associations between temperature, pulse rate, blood pressure (BP), and respiration rate in COVID-19 and flu diagnosed patients. Data from 1,363 COVID-19 (March 3, 2020, to February 27, 2021) and 507 flu (October 1, 2017, to September 30, 2018) diagnosed patients with reported demographic dimensions (age, first race, and sex) and office visit dimensions (BMI, diastolic BP, pulse rate, respiration rate, systolic BP, and temperature) was obtained, including day of diagnosis and additional encounter visits 60 days before and after first unique diagnosis. Patients with COVID-19 or flu were disproportionately obese, with 93% of COVID-19 and 79% of flu patients with BMI ≥ 30. Most striking, Black women 50–64 years of age disproportionately carried the burden of disease. At the time of diagnosis, temperature was significantly increased for all patients, yet pulse rate was only significantly increased for flu diagnosis, and BP was not significantly different in either. Our findings show the need for more complete demographic and office visit dimension data from patients during epidemic and pandemic events and support further studies needed to understand association between vital signs and predicting respiratory disease.

Keywords COVID-19 · Flu · Mississippi · Pulse rate · Temperature · Racial disparities

Introduction

Racial and ethnic disparities are clearly evident in certain types of infections common among women [1, 2]. These include sexually transmitted diseases and urinary tract infections, and more recently, women seem to be disproportionately positive for SARS-CoV-2, the virus that causes COVID-19 disease [3–5]. The Black population in Mississippi (MS) is 39% of the total population, the highest proportion in the USA. As of April 2021, the Black population in MS represented only one-third of COVID-19 cases, but over half of associated deaths [6]. Racial statistics are limited for other diseases, such as seasonal influenza virus due to being sparsely reported in 14 states [7]. Risk factors associated with increased morbidity and mortality for COVID-19 and other respiratory infections including influenza include a body mass index (BMI) of 30 or greater and advanced age [8–16]. Given that Mississippi has the highest prevalence of obesity (41%; obesity defined as BMI ≥ 30 [17]) in the USA, along with 46% of the Black population being obese or morbidly obese, the potential impact of respiratory disease is of concern. Additionally, sex differences in disease outcome are now coming to light, especially in COVID-19, indicating a larger risk of morbidity for men [18].

The inflammatory response during an infection can induce a fever that leads to an elevated heart rate [19]. For COVID-19, while there have been some studies to examine heart rate, none has identified a significant association with increased heart rate [20]. In contrast, other studies using wearable data monitoring devices that measure heart rate,
activity, and sleep have identified associations with influenza [21]. Early detection using these types of measurement have the potential to reduce risk of morbidity, mortality, and disease transmission through earlier intervention strategies. Thus, distinguishing patterns among vital signs, body temperature, pulse rate, respiration rate, and blood pressure, can help to more rapidly and accurately diagnose a potential respiratory virus.

The University of Mississippi Medical Center (UMMC) is Mississippi’s only academic medical center and treats patients from across the state through its main campus, located in Jackson as well as two community hospitals. The objective of this study was to utilize UMMC’s extensive clinical database to investigate the potential association between temperature, pulse rate, blood pressure, and respiration rate with COVID-19 and flu. In particular, the large Black population in Mississippi provides an opportunity to examine whether racial differences exist between these physiological parameters and respiratory diseases.

Materials and Methods

Data extracted from the UMMC Patient Cohort Explorer (PCE) [22] contains patient data derived from the electronic health record used on-site (EPIC) from January 1, 2013, through February 28, 2021. The data warehouse includes de-identified and date-shifted clinical information and does not include any protected health information (PHI). Pursuant to 45 CFR 46, use of this database does not meet the definition of human subjects’ research and does not require IRB review.

COVID-19 patients were identified through the PCE COVID-19 Research Registry dashboard (March 3, 2020, to February 27, 2021), which were diagnosed as SARS-CoV-2 positive either by rapid antigen assay, reverse transcriptase polymerase chain reaction, or seropositivity. Patients with a diagnosis of flu were identified by the encounter diagnosis name “influenza A” or “influenza B,” determined by rapid antigen assay, reverse transcriptase polymerase chain reaction, direct or indirect fluorescent staining, or viral culture. To note, diagnoses are only defined by a positive test and may not reflect a symptomatic state. The total number of flu patients (both influenza A and influenza B) from the 2017–2018 season is included in this study, October 1, 2017, to September 30, 2018. To control for variability for comorbidities, flu diagnosed patients without any other previous diagnoses are compared to those previously diagnosed with essential (primary) hypertension (ICD-10 I10 [23]).

Mississippi population estimates of age, sex [24], race [25], and BMI [26] establish per capita frequencies. Chi-squared test (α = 0.05) evaluates the association of positivity cases for disease diagnosis, age, sex, race, and BMI. One-way ANOVA (α = 0.05) assesses differences in office visit dimensions (BMI, diastolic blood pressure (BP), pulse rate, respiration rate, systolic BP, and temperature) among time relative to diagnosis, with post hoc tests Dunnett’s multiple comparisons test (α = 0.05) for differences between day of diagnosis and other time periods and Sidak’s multiple comparisons test for differences among data within a time period.

Results

In total, 24,030 patients were identified as SARS-CoV-2 positive (COVID-19 diagnosis) from March 3, 2020, to February 27, 2021, and 4,050 patients were influenza A or influenza B positive (flu diagnosis) from October 1, 2017, to September 30, 2018 (Table S1). Patients with reported demographic dimensions (age, race, and sex) and office visit dimensions (BMI, diastolic blood pressure (BP), pulse rate, respiration rate, systolic BP, and temperature) for day of diagnosis and additional encounter visits within 60 days before and after first unique viral disease diagnosis are included in the analysis; 1,363 COVID-19 diagnosed patients and 507 flu diagnosed patients are ultimately included in the analysis (Table S2).

General Patient Demographics for COVID-19 and Flu

Patients diagnosed with COVID-19 or flu are stratified based on age or BMI (Fig. 1). During the time period studied, the peak of COVID-19 diagnosis occurs in 50–64-year-olds, which is approximately double the number of patients at the tail ends of the age spectrum. In contrast, there is no apparent association between flu diagnosis and age, as diagnosis was similar across all the age groups. Patients with COVID-19 or flu diagnoses are disproportionately obese, with 93% of COVID-19 and 79% of flu patients having a BMI ≥ 30.

The diagnoses grouped by race and sex reveal that more Black females have a diagnosis of COVID-19 or flu compared to White females, while Black males have a diagnosis of COVID-19 twice that compared to white males (Fig. 2A and C). Black female patients ages 50–64 disproportionately represent infected patients in the Black population, with 18% COVID-19 diagnosis and 13% flu diagnosis. Additionally, 70% of COVID-19 and 53% of flu patients are Black with a BMI ≥ 30 and 60% of COVID-19 and 54% of flu patients are females with BMI ≥ 30 (Fig. 2B and D). Black female patients have the most diagnosis for COVID-19 and have BMI > 40 (22%) and for flu have BMI between 30 and 39 (17%).

Overall, there are disproportional diagnoses in Black female patients composing the highest number of COVID-19 (51%) and flu (45%) patients. Additionally,
Black female patients with BMI $\geq 30$ are the majority infected with both COVID-19 (42%) and flu (30%). Black female patients with the highest number of diagnoses are 50–64 years old with BMI $\geq 30$ with 15% of
total COVID-19 and 12% of total flu patients being Black female of 50–64 years old with BMI ≥ 30.

**Disease Dynamics Based on Vital Signs**

The data of vital sign measurements are presented relative to first unique viral diagnosis and grouped into −60 to −31 days before diagnosis (−2 months), −30 to −1 days before diagnosis (1 month), day of diagnosis (0), 1 to 30 days after diagnosis (1 month), and 31 to 60 days after diagnosis (2 months). For temperature in COVID-19 patients, there is a significant difference among the time points \( p < 0.0001 \); one-way ANOVA \( (\alpha = 0.05) \) with day of diagnosis significantly different from every other time point \( (vs -2\text{ months} \ p < 0.0001 \text{ and vs 1 month and 2 months} \ p = 0.0001 \text{ and 0.0002, respectively}; \text{ Dunnett’s multiple comparisons test (} \alpha = 0.05; \text{ Fig. 3a}) \). COVID-19 patients have an average increase of 0.54°F from the average temperature throughout the timeline; pre- and post-diagnosis temperature range from 93.2 to 103.1°F and day of diagnosis temperatures range from 96.0 to 101.6°F (Fig. 3b). Temperature is also significantly different for flu patients, \( p < 0.0001 \), with the day of diagnosis having statistical difference between every other time point \( (p < 0.0001 \text{ for all}; \text{ Fig. 3c}) \). Flu patients have an average increase of 1.00°F from the average temperature throughout the timeline; pre- and post-diagnosis temperature range from 95.1 to 100.7°F and day of diagnosis temperatures range from 96.1 to 103.0°F (Fig. 3d).

COVID-19 diagnosed patients do not have significant differences in pulse rate across the time of study. For flu patients, there is a significant difference within the timeline \( (p < 0.0001) \) with day of diagnosis being significantly different from every time point \( (p < 0.0001 \text{ for all}; \text{ Fig. 4a}) \). Flu patients have an average increase of 7.6 beats per minute (bpm) from the average pulse rate throughout the timeline; pre- and post-diagnosis pulse rate range from 51.0 to 146.4 bpm and day of diagnosis pulse rate ranges from 53.0 to 164.0 bpm (Fig. 4b).

**Demographic Biases in Diagnosis Dynamics**

Given the potential impact of patient demographics on diagnosis as described above, we examined clinical parameters stratified by sex and race at each time point. This analysis was focused on flu patients due to more office visit dimension data available for patients before, during, and after flu diagnosis.

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**Fig. 3** Temperature dynamics during COVID-19 and flu diagnoses. Temperature for each patient was grouped relative to day of diagnosis −60 to −31 days before diagnosis (−2 months), −30 to −1 days before diagnosis (1 month), day of diagnosis (0), 1 to 30 days after diagnosis (1 month), and 31 to 60 days after diagnosis (2 months) for COVID-19 (A) and flu (C) diagnosed patients; mean denoted by cross (* \( p\text{-value}<0.0001 \)). The average temperature for each day within −60 to 60 days post diagnosis (grey circles) and 7-day moving average (Black line) shows the time course of temperature dynamics in COVID-19 (B) and flu (D) diagnosed patients.
The average and standard deviation of the office visit dimensions were calculated for each of the discrete time points for four sex and race groups (Black female, White female, Black male, and White male) and tested for statistical difference using one-way ANOVA independently (Table 1). For temperature, all four groups are statistically denoted by cross (* p-value < 0.0001). The average temperature for each day within −0 to 60 days post diagnosis (gray circles) and 7-day moving average (black line) show the time course of pulse rate dynamics in flu diagnosed patients (B).

Table 1 Average and standard deviation (in parenthesis) are reported for each office visit dimension for each sex and race group for the discrete time periods of −60 to −31 days before diagnosis (−2 months), −30 to −1 days before diagnosis (1 month), day of diagnosis (0), 1 to 30 days after diagnosis (1 month), and 31 to 60 days after diagnosis (2 months). One-way ANOVA (α = 0.05) was performed for each office visit dimension to obtain p-value.

|                          | −2 months | −1 month | 0       | 1 month | 2 months | p-value |
|--------------------------|-----------|----------|---------|---------|----------|---------|
| Female Black or African American (n = 225) | | | | | | |
| Temperature (F)          | 98.02 (0.58) | 98.05 (0.64) | 99.36 (1.19) | 98.07 (0.73) | 98 (0.52) | < 0.0001 |
| Pulse rate               | 81.98 (11.67) | 83.71 (14.86) | 94.74 (14.88) | 83.12 (12.45) | 82.38 (12.63) | < 0.0001 |
| Systolic BP              | 134.7 (17.61) | 134.1 (25.49) | 134.5 (19.49) | 129 (19.94) | 131.3 (21.58) | 0.2234 |
| Diastolic BP             | 78.33 (11.47) | 78.27 (13.61) | 80.44 (13.23) | 77.02 (11.92) | 75.94 (12.05) | 0.1302 |
| Respiration rate         | 18.54 (2.15) | 17.83 (1.99) | 17.74 (1.94) | 17.8 (2.1) | 18.25 (1.81) | 0.1756 |
| Female White or Caucasian (n = 104) | | | | | | |
| Temperature (F)          | 97.93 (0.67) | 97.92 (0.84) | 98.86 (1.26) | 97.95 (0.63) | 97.9 (0.5) | < 0.0001 |
| Pulse rate               | 79.2 (13.75) | 82.44 (13.67) | 96.4 (18.61) | 81.84 (13.18) | 79.93 (14.66) | < 0.0001 |
| Systolic BP              | 130 (19.48) | 129.7 (22.37) | 127.6 (13.46) | 129.2 (12.67) | 131.2 (16.08) | 0.3626 |
| Diastolic BP             | 73.36 (12.9) | 73 (12.36) | 76.25 (9.71) | 72.72 (12.83) | 75.18 (10.75) | 0.4004 |
| Respiration rate         | 17.95 (1.76) | 18.5 (2.26) | 18.2 (1.53) | 19.05 (2.55) | 18.83 (1.65) | 0.279 |
| Male Black or African American (n = 111) | | | | | | |
| Temperature (F)          | 97.89 (0.67) | 98.9 (1.13) | 97.99 (0.83) | 97.99 (0.83) | 97.83 (0.68) | < 0.0001 |
| Pulse rate               | 86.91 (15.99) | 89.95 (18.4) | 88.61 (15.48) | 90.75 (17.17) | 87.52 (12.46) | 0.7405 |
| Systolic BP              | 125.8 (16.61) | 131.6 (16.49) | 132.9 (19.83) | 129 (17.59) | 130.1 (15.99) | 0.3316 |
| Diastolic BP             | 76.75 (8.67) | 77.66 (11.99) | 80.25 (12.65) | 80.45 (10.47) | 79.58 (11.77) | 0.4029 |
| Respiration rate         | 18.38 (1.26) | 18.59 (2.16) | 18.33 (1.98) | 18.33 (1.98) | 18.46 (1.61) | 0.9796 |
| Male White or Caucasian (n = 67) | | | | | | |
| Temperature (F)          | 97.76 (0.73) | 97.96 (0.81) | 98.82 (1) | 97.86 (0.69) | 98.11 (0.52) | < 0.0001 |
| Pulse rate               | 73.52 (40) | 79.52 (40) | 85.8 (40) | 82.54 (40) | 78.67 (40) | 0.0546 |
| Systolic BP              | 131.9 (12.01) | 126.4 (15.43) | 127.2 (19.9) | 133.5 (16.53) | 126.7 (17.15) | 0.3894 |
| Diastolic BP             | 76.37 (9.11) | 75 (11.86) | 77.61 (12.35) | 78.35 (13.02) | 71.11 (10.11) | 0.0942 |
| Respiration rate         | 17.84 (2) | 18 (1.71) | 18.84 (1.99) | 19.15 (1.14) | 18 (2.37) | 0.2686 |
significant (p < 0.0001, for all). Comparison among the
groups within a time point, using Sidak’s multiple compar-
isons test (α = 0.05), shows that White males have a signifi-
cantly higher temperature at −1 month before flu diagnosis
than all other groups. At day of diagnosis, White females
and white males are not significantly different from each
other, yet Black females and Black males are significantly
different from all other groups.

For pulse rate, only the female groups have a signifi-
cant difference in pulse rate (p < 0.0001) while the male
groups are not significantly different (Fig. 5). Black females
and White females have an average increase of 12.0 and
15.6 bpm, respectively, from the average pulse rate through-
out the timeline; pre- and post-diagnosis pulse rates range
from 55.0 to 130.0 bpm (Black females) and 54.0 to
116.0 bpm (White females) and day of diagnosis pulse rates
range from 72.0 to 164.0 bpm (Black females) and 69.0 to
145.0 bpm (White females). Additionally, comparison
among the groups within a time point shows differences in
race and/or sex at −2 months, day of diagnosis, and 1 month
(Fig. 5). Overall, White females have a greater increase in
pulse rate at day of diagnosis while Black females have a
larger range or increased pulse rates at day of diagnosis.
Blood pressures and respiration rate were not significantly
different at day of diagnosis for any of the groups.

To determine if pre-existing diseases could bias office
visit dimensions, patients with no pre-existing diagnosis
(n = 57) were compared to those with essential (primary)
hypertension (n = 234). There is a significant difference
in systolic BP at day of diagnosis, with CVD having an
average systolic BP of 131.29 compared to 138.90 of flu
only patients. Temperature is significantly lower in CVD
patients −1 month and day of diagnosis compared to flu only
patients; the difference in temperature increase from average
across the timeline for CVD patients was 1.10°F while flu
only patients had an average increase of 1.13°F. Pulse rate
significantly changed across visits, peaking at diagnosis, but
was not significantly different between those with flu only
compared to flu and CVD (Table S3). The average increase
in pulse rate for CVD patients was 11.42, slightly higher
than flu only patients which was 8.30. Thus, we found no
significant evidence that pre-existing diseases affect office
dimensions.

Discussion

For Mississippi, COVID-19 cases and deaths are reported
by race but not for flu virus at either the country (CDC) or
state (MSDH) levels. Age, sex, and race seem to play a vital
role in diagnosis population statistics, and more complete
demographic reports are necessary to understand the role of
disparities. Factors, such as reported BMI, show a vital role
in COVID-19 and flu patients. Obesity was a risk factor for
severe morbidity or mortality upon infection for COVID-19
[10] and the 2009 pandemic H1N1 virus [27–33] as well
as an increasing risk of hospitalization for both pandemic
and non-pandemic flu infections with an increase in BMI
[34, 35].

Global data shows little difference of COVID-19 case
numbers based on sex [36], yet in this study (during the
period studied), we found that women in Mississippi have
the highest number of COVID-19 and flu diagnoses. Tak-
ing into account sex disparities across racial groups, we
found overall that Black women have the greatest number
of diagnoses for both viruses further showing the limitation
of unidimensional reporting that relies on only sex or racial
statistics alone (also highlighted by [37]). This study adds
another example of how Black women in the US continue

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**Fig. 5** Pulse rate dynamics
during flu diagnosis for 4 sex
and race groups. Pulse rate
for each patient was grouped
relative to day of diagnosis −60
to −31 days before diagnosis
(−2 months), −30 to −1 days
before diagnosis (1 month), day
of diagnosis (0), 1 to 30 days
after diagnosis (1 month), and
31 to 60 days after diagnosis
(2 months); mean denoted by
cross. Comparison among the
groups within a time point was
completed with Sidak’s multiple
comparisons test (α = 0.05), *
p-value < 0.05
to have health inequalities that contribute to greater burden of disease [38].

Social distancing has been recommended in past pandemics, including influenza virus, to limit or stop the spread of highly transmissible diseases [39, 40]. Those groups with less social isolation and social distancing may be more likely to be infected with highly transmissible viruses. This could be the case with middle-aged women in this study with the most diagnosis. Higher social interactions, deferring from CDC guidelines for slowing the spread of COVID-19 [41], and coming into contact with children and teenagers have all been noted to play an important role in disease transmission [42]. In all data, for COVID-19, there were 190 patients diagnosed with pregnancy or postpartum with age ranging from 14–42 and for flu, there were 114 patients diagnosed with pregnancy or postpartum in the 18–38 age range. However, all these patients were excluded from this study due to lack of office visit dimensions. Additionally, with the majority of diagnosis being infected women ranging from 50 to 64 years old age group, one factor that is not addressed in the PCE database is if these women are currently in menopause. Menopause, typically occurring between ages of 50 and 52, has been linked, albeit in lean women, lower lung function in menopausal women [43]. Although there is no racial difference in the start of menopause, there is a significant difference in location with those in southern states which have earlier menopause than the rest of the USA [44].

The overarching goal for the identification of vital signs using non-invasive monitoring is to be able to determine a viral infection before symptom onset. Asymptomatic infections are common in COVID-19 and flu, and transmission still occurs to naïve individuals [45–47]. Faster application of interventions can reduce virus replication and spread; for example, the commonly used anti-flu drug oseltamivir (Tamiflu) is more effective when taken within 48 h of symptom onset [48], yet most do not seek clinical intervention until 3–8 days post symptom onset [49].

In this study, we primarily saw significant variations from the average at the day of disease diagnosis but not before. We show a significant difference from baseline in pulse rate of flu diagnosed patients, similar to other findings [50], but no significant difference in COVID-19 diagnosed patients as others were unable to find as well [20]. On the other hand, increased temperature is a well-known marker for infection. A fever is defined as a temperature at or above 100.4°F, yet the average oral temperature has decreased from 98.6 to 97.9°F without a change in guidelines for fever. As found here, an increase of 0.5°F for COVID-19 and 1.0°F for flu could be substantial enough to suspect a viral infection.

We do not see a significant change in blood pressure before and after diagnosis from date of diagnosis. This could be due to multiple factors including direct and indirect interventions, including cold medicines and blood pressure medicines, respectively, that could potentially stabilize blood pressure from increasing during infection. Those with CVD and flu diagnosis do have higher BP than those with flu diagnosis only, yet there is still no difference in BP on day of diagnosis compared to their BP before and after diagnosis.

Importance of disease outcome, including mortality, could further inform about differences found at day of diagnosis, leading to information for prognosis prediction. One major caveat to consider in this study relates to lack of patient death information. Neither date of death nor cause of death is logged; additionally, death is only added into EPIC if mortality is reported to the hospital, or the patient has died within the hospital umbrella system. With this in mind, we are unable to assess risk of disease fatality in this study and their relation to difference found at day of diagnosis in order to compare to national averages for COVID-19 and flu found by others [51–53].

Conclusions

Our findings show the need for more complete demographic and office visit dimension data from patients during epidemic and pandemic events. In this study, we find that disease burden is greater in Black women in 50–64 age group for COVID-19 and flu virus in the Mississippi population. For all patients at the time of diagnosis, temperature is significantly increased for both respiratory viruses and pulse rate is significantly increased for flu virus diagnosis. This data is consistent with previous research and supports the idea that these vital sign measurements could provide early indication of infection with application of wearable devices.

Supplementary Information  The online version contains supplementary material available at https://doi.org/10.1007/s40615-021-01213-2.

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Declarations

Conflict of Interest  The authors declare no competing interests.

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