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Clinical outcomes of COVID-19 in patients with sickle cell disease and sickle cell trait: A critical appraisal of the literature

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ABSTRACT

Individuals with sickle cell disease (SCD) and sickle cell trait (SCT) have many risk factors that could make them more susceptible to COVID-19 critical illness and death compared to the general population. With a growing body of literature in this field, a comprehensive review is needed. We reviewed 71 COVID-19-related studies conducted in 15 countries and published between January 1, 2020, and October 15, 2021, including a combined total of over 2000 patients with SCD and nearly 2000 patients with SCT. Adults with SCD typically have a mild to moderate COVID-19 disease course, but also a 2- to 7-fold increased risk of COVID-19-related hospitalization and a 1.2-fold increased risk of COVID-19-related death as compared to adults without SCD, but not compared to controls with similar comorbidities and end-organ damage. There is some evidence that persons with SCT have increased risk of COVID-19-related hospitalization and death although more studies with risk-stratification and properly matched controls are needed to confirm these findings. While the literature suggests that most children with SCD and COVID-19 have mild disease and low risk of death, some children with SCD, especially those with SCD-related comorbidities, are more likely to be hospitalized and require escalated care than children without SCD. However, children with SCD are less likely to experience COVID-19-related severe illness and death compared to adults with or without SCD. SCD-directed therapies such as transfusion and hydroxyurea may be associated with better COVID-19 outcomes, but prospective studies are needed for confirmation. While some studies have reported favorable short-term outcomes for COVID-19 patients with SCD and SCT, the long-term effects of SARS-CoV-2 infection are unknown and may affect individuals with SCD and SCT differently from the general population. Important focus areas for future research should include multi-center studies with larger sample sizes, assessment of hemoglobin genotype and SCD-modifying therapies on COVID-19 outcomes, inclusion of case-matched controls that account for the unique sample characteristics of SCD and SCT populations, and longitudinal assessment of post-COVID-19 symptoms.

1. Introduction

Coronavirus disease 2019 (COVID-19) [1–3] is an infectious disease caused by a newly discovered coronavirus (SARS-CoV-2), which has impacted millions of people around the world and caused unprecedented challenges to healthcare systems. Severe COVID-19 clinical manifestations include acute respiratory distress syndrome (ARDS), systemic inflammation, sepsis, multi-organ failure, and death, among others [1–3]. Survivors of SARS-CoV-2 infection may experience lingering symptoms and long-term health problems [4].

Sickle cell disease (SCD) is an inherited red blood cell disorder caused by a single amino acid genetic mutation in the beta-chain of the human hemoglobin protein [5,6]. Patients with SCD experience chronic hemolytic anemia, recurrent vascular occlusion, insidious vital organ

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deterioration, early mortality, and poor quality of life [5,6]. Repeated vaso-occlusive events may result in functional asplenia and immune deficiency in early childhood leading to life-long increased susceptibility to serious bacterial infections. Acute and chronic pain and other serious health conditions such as acute chest syndrome (ACS) and stroke [7] are also common in patients with SCD. SCD predominantly affects Black individuals. In the United States, there are about 100,000 people living with SCD, with 1 in 500 African Americans being affected by the disease [8].

Individuals with sickle cell trait (SCT) may also exhibit adverse health effects including exertional rhabdomyolysis, pulmonary embolism, splenic infarction, and renal damage [9,10]. Approximately 300 million people worldwide and 1 to 3 million in the United States (8% to 10% of African Americans) live with SCT [10–12].

There is conflicting evidence whether patients with SCD or SCT experience more severe COVID-19 disease, with higher morbidity and mortality rate than the general population. SCD pathophysiology results in chronic anemia, endothelial dysfunction, chronic inflammation, immunocompromised status, and hypercoagulability, all of which have been identified as risk factors for worse COVID-19 outcomes [13–16]. Individuals with SCT do have a hypercoagulable state and sickling pathophysiology in hypoxic environments such as the renal medulla and could potentially be at higher risk for severe COVID-19 outcomes as well.

In this paper, we present a comprehensive review of the literature of COVID-19 patients with SCD and SCT and highlight both the conclusions and limitations drawn from the literature, which can serve as important considerations for future research. We also discuss clinical course, hospitalization trends, mortality, risk factors, and SCD-disease modifying therapies among children and adults (separately) with SCD and SCT (separately) who have COVID-19 disease.

2. Methods

A structured literature search was conducted using PubMed to include all relevant articles that were dated between January 1, 2020, and October 15, 2021. The selection process is detailed in Fig. 1. The review was conducted according to the PRISMA 2020 27-item checklist [17]. Titles and abstracts were searched to identify the literature related to COVID-19 and SCD/SCT. The primary search term used was ‘COVID-19 sickle cell’ which yielded 157 articles. Additional references were retrieved from the bibliographies of the articles identified by the primary search. Of the original 164 records screened, 50 were excluded because they were unrelated to SCD and COVID-19, and 43 were excluded because they were not available in English or did not include original clinical data on SCD or SCT patients with positive or suspected COVID-19. After this filter, a total of 71 publications were retained and systematically reviewed.
2.1. Statistical analysis

In this systematic review, we used descriptive statistics to summarize aggregated demographic and clinical data of SCD and SCT patients with SARS-CoV-2 infection based on the literature included in this review (Table 1), including median and interquartile range for age, vitals, and laboratory values. The categorical variables female sex, hemoglobin genotype, and comorbidities were described as n (% of the sample). The overall mortality rate for SCD/non-SCD was calculated as the total number of COVID-19-related deaths among SCD/non-SCD patients reported in observational cohort studies divided by the total number of SCD/non-SCD patients with COVID-19 reported by the same studies. Relative risk of death for SCD was estimated as the absolute risk in the SCD group divided by the absolute risk in the non-SCD group.

3. Patients with COVID-19 and sickle cell disease

Of the 71 reports identified in this review, 67 studies (25 observational cohort studies and 42 case studies) reported on SCD patients describing a combined total of 2290 patients with SCD and COVID-19. A detailed summary of study parameters and main study findings are presented in Table 2 (observational studies) and Table 3 (case studies).

3.1. Sample characteristics

There were 2290 persons with SCD and SARS-CoV-2 infection analyzed by all studies combined and included in this review (Table 1). The SCD cohort had a median age of 25 years (interquartile range = 17–39 years), was 56% female, and 62% had HbSS hemoglobin genotype (Table 1). The relatively young age of patients with SCD and COVID-19 could be explained by the short life expectancy of SCD patients (43 years in 2017 [18]) relative to the general population (79 years in 2017 [19]). Despite young age, SCD patients had substantial comorbidities (i.e., hypertension, asthma/COPD, and chronic kidney disease), consistent with end-organ damage from SCD [20,21].

At hospital presentation, oxygen saturation was 95%, body temperature was 101.6°F, and BMI was 28. Laboratory values were above normal range for lactate dehydrogenase, bilirubin, white blood cell count, C-reactive protein, and D-dimer; below normal range for hemoglobin; and within normal range for lymphocyte count. These altered laboratory values are consistent with SCD-related manifestations of red blood cell dysfunction [6,22]. Elevated lactate dehydrogenase and D-dimer could be suggestive of more severe COVID-19 disease [23] but could also result from intravascular hemolysis, ischemia-reperfusion damage and tissue necrosis associated with SCD and could be further elevated during acute vaso-occlusive crisis [24].

3.2. Clinical course

Adult SCD patients typically experience mild to moderate COVID-19-related symptoms [25–32], which is characterized by few respiratory symptoms and not requiring intensive care unit (ICU) admission. Large US registry studies found that approximately 80% of adult SCD patients experience mild to moderate symptoms (or no symptoms at all) [29,30] and that the type of COVID-19-related symptoms, including cough/fever or more severe symptoms such as respiratory failure, were not different from patients without SCD [33]. However, SCD-related comorbidities, in particular acute pain has been found highly prevalent during COVID-19, and history of pain was associated with increased risk of hospitalization [29].

A few smaller cohort studies observed more severe COVID-19 disease course among adult SCD patients [34–37]. Severe disease course is typically characterized by patients needing escalated care and prolonged hospitalization due to severe COVID-19 symptoms (e.g., hypoxia, pneumonia, prolonged fever, multisystem inflammation) or severe SCD-related complications (e.g., ACS, severe pain crisis), or both. One study observed increased hospitalization, mild to moderate pneumonia, and intubation among SCD patients as compared to age-matched healthcare professionals [37]. Another study identified older patients, pre-existing conditions, and end-organ damage as high risk factors for severe COVID-19 disease course and poor outcome [36].

3.3. Hospitalization trends

There is strong evidence that patients with SCD have an increased risk of COVID-19-related hospital admissions. Large cohort studies in the UK and US have reported a 2- to 7-fold increased risk of COVID-19-related hospital admission for patients with SCD relative to the general population [33,38,39]. Most observational studies reported hospitalization rates of greater than or equal to 40% among patients with SCD and SARS-CoV-2 infection [26,28,30,32,35,36,33,39,40], which remained increased after age, gender, and race adjustment when compared to the general population [33,38,39]. The hospitalization rate of individuals with SARS-CoV-2 infection in the general population is markedly lower. For example, peak hospitalization for the US population was 20.5% (first week of January 2021; https://covid.cdc.gov/covid-data-tracker/) and 19.2% in Black patients without SCD in a large US registry study [33]. Case studies show even higher hospitalization trends, although this may be in part due to patient selection bias. Of the 9 case studies that reported both hospitalized and non-hospitalized patients, the COVID-19-related hospitalization rate for persons with SCD was 72% [44,61] [41–45].

There could be several explanations for the high hospitalization rate observed among patients with SCD. We hypothesize that the high baseline hospital admission and readmission rate for patients with SCD, independent of COVID-19, may have been contributing factors, in
Table 2

Observational cohort studies of patients with SCD and COVID-19. Authors listed in alphabetical order.

| Authors                  | Country       | Sample size (n M/F) | Age (years) | Hb genotype (n) | SCD therapy (n) | Control group | Main findings                                                                 |
|--------------------------|---------------|--------------------|-------------|-----------------|-----------------|---------------|-------------------------------------------------------------------------------|
| AbdulRahman et al. [25]  | Bahrain       | 38 (11/27)         | 36          | n/a             | n/a             | COVID-19 w/o SCD | SC is not a risk factor for worse COVID-19 outcomes in hospitalized pts. compared to non-SCD pts. hospitalized with COVID-19, 14 of 31 SCD pts. with COVID-19 required admission to the ICU. Hosp. rate n (%): 1 (2.6) Mort. rate n (%): 58 (3.3) |
| Alhumaid et al. [34]    | Saudi Arabia  | 31 (n/a)           | n/a         | n/a             | n/a             | None           | None                                                                                                                                   |
| Alkindi et al. [26]     | Oman          | 50 (27/23)         | 31          | HbSS (10), HbSβ (29), Unknown (11) | BT (26)         | SCD w/o COVID-19 | Although COVID-19 may trigger VOC onset, it does not increase mortality compared to non-COVID-19 pts. with VOC. Hosp. rate n (%): 34 (68) Mort. rate n (%): 47 (70) |
| Al Yazidi et al. [64]   | Oman          | 7 (n/a)            | n/a         | n/a             | BT (2)          | None           | SCD is the most common comorbidity associated with COVID-19 pediatric admission. Children with SCD and COVID-19 are more susceptible to complications. SC pts. did not have increased risk of morbidity or mortality; VOC complicates infection; Older pts. are at higher risk and should be monitored carefully. Hosp. rate n (%): 4 (67) Mort. rate n (%): 2 (2) |
| Arlet et al. [27]       | France        | 83 (38/45)         | 30          | HbSS or HbSβ (71), HbSC (8), HbSβ+ (4) | HU (38), BT (31) | COVID-19 w/o SCD | SC pts. did not have increased risk of morbidity or mortality; VOC complicates infection; Older pts. are at higher risk and should be monitored carefully. Hosp. rate n (%): 4 (67) Mort. rate n (%): 2 (2) |
| Balanchivadze et al. [28] | USA          | 6 (3/3)            | 38          | HbSS (4), HbSC (1), HbSβ+ (1) | BT (2)          | None           | SC pts. generally had mild disease course with lower chances of intubation, ICU admission, and death. Hosp. rate n (%): 4 (67) Mort. rate n (%): 2 (2) |
| Boga et al. [37]        | Turkey        | 39 (17/22)         | 35          | HbSS (23), HbSβ+ (15), HbSE (1) | HU (25), BT (8) | COVID-19 w/o SCD | SC pts. had more severe disease course (pneumonia, hosp., intubation) than non-SCD healthcare professionals. Among seropositive patients, none had displayed symptoms except one who was hospitalized for mild vaso-occlusive symptoms with a favorable outcome. Hosp. rate n (%): 10 (26) Mort. rate n (%): 9 (7) |
| Brousse et al. [70]     | France        | 39 (n/a)           | 12          | HbSS (35), HbSC (3), HbSβ+ (1) | HU (25)         | SCD w/o COVID-19 | Among seropositive patients, none had displayed symptoms except one who was hospitalized for mild vaso-occlusive symptoms with a favorable outcome. Hosp. rate n (%): 1 (2.6) Mort. rate n (%): 0 (0) |
| Clift et al. [38]       | UK            | 287 (n/a)          | n/a         | n/a             | n/a             | COVID-19 w/o SCD | SC pts. had 4-fold and 2.6-fold increased risk of hospitalization and death due to COVID-19, respectively. Hosp. rate n (%): 40 (14) Mort. rate n (%): 23,561 (4.4) |
| De Sanctis et al. [35]  | Oman          | 3 (1/2)            | 27          | n/a             | HU (3), BT (1)  | None           | SC and other comorbidities can aggravate COVID-19 severity despite favorable outcome; Two SCD pts. had worsening anemia; Respiratory complications rapidly improved after red blood cell exchange BT in one patient. Hosp. rate n (%): 2 (67) Mort. rate n (%): 0 (0) |
| Fisler et al. [90]      | USA           | 2 (n/a)            | n/a         | n/a             | n/a             | None           | Two SCD pts. were admitted to ICU for ACS. Hosp. rate n (%): n/a Mort. rate n (%): n/a |
| Gampel et al. [65]      | USA           | 1 (1/0)            | 12          | HbSC            | n/a             | None           | SC pts. with h/o prior pulmonary and cardiac complications died of COVID-19 disease. Hosp. rate n (%): n/a Mort. rate n (%): 1 (100) |
| Hippisley-Cox et al. [52] | UK           | n/a               | n/a         | n/a             | n/a             | COVID-19 w/o SCD | SC pts. vaccinated had 7.7-fold increased risk of COVID-19 mortality. Hosp. rate n (%): n/a Mort. rate n (%): n/a |
| Hoogenboom et al. [39]  | USA           | 53 (28/25)         | 30          | HbSS (39), HbSC (11), HbSβ (3) | n/a             | Matched and unmatched COVID-19 pts. w/o SCD | SC pts. were more likely to visit the ED and be hospitalized than the general population. Mortality, severe disease, and other outcomes in SCD pts. was not different from controls. Hosp. rate n (%): 39 (74) Mort. rate n (%): 3 (8) |

(continued on next page)
| Authors          | Country | Sample size (M/F) | Age (years) | HB genotype (n) | SCD therapy (n) | Control group | Main findings                                                                 | SCID Hosp. rate n (%) | SCID Mort. rate n (%) | Non-SCID Hosp. rate n (%) | Non-SCID Mort. rate n (%) |
|-----------------|---------|-------------------|-------------|-----------------|-----------------|---------------|-----------------------------------------------------------------------------|----------------------|----------------------|------------------------|------------------------|
| Kamdar et al. [40] | USA     | 30 (n/a)          | n/a         | n/a             | HU (19), BT (3)  | None          | SCD pts. had more ED visits and hospitalization than other hematological pts. with COVID-19. | 14 (47)              | 0 (0)                | n/a                    | n/a                    |
| Konte et al. [100] | Netherlands | 5 (n/a)       | n/a         | n/a             | None            | None          | Low incidence of COVID-19 in SCD pts. presenting with VOC, suggesting that COVID-19 is not a major provoking factor for VOC. SCD pts. over 50yo with preexisting conditions with elevated creatinine, LDH, and D-dimer are at higher risk of death regardless of genotype or sex. All deaths occurred in pts. not on HU. | n/a                  | n/a                  | n/a                    | n/a                    |
| Minniti et al. [36] | USA     | 66 (30/36)        | 34          | HbSS or HbSβ+ (47), HbSC (14), HbSβ+ (5) | HU (28), BT (30) | None          | SCD pts. had relatively mild disease course. Black SCD pts. are more likely to be hospitalized and develop pneumonia and pain than Black pts. w/o SCD but do not differ in mortality. | 50 (76)              | 7 (11)               | n/a                    | n/a                    |
| Mucalo et al. [29] | USA     | 364 (187/176)     | 11          | HbSS or HbSβ+ (283), HbSC or HbSβ+ (98) | HU (203), BT (39) | None          | SCD children with h/o pain, renal, and heart/lung comorbidities are at higher risk of worse COVID-19 outcomes. | 146 (40.1)           | 1 (0.3)               | n/a                    | n/a                    |
| Mucalo et al. [29] | USA     | 386 (159/220)     | 31          | HbSS or HbSβ+ (261), HbSC or HbSβ+ (113) | HU (191), BT (53) | None          | SCD Adults with h/o pain are at higher risk of worse COVID-19 outcomes. | 231 (59.8)           | 18 (4.7)              | n/a                    | n/a                    |
| Nathan et al. [60] | India   | 178 (75/101)      | 17          | n/a             | BT (2)          | None          | Two SCD pts. developed ACS and required ICU and NIV. All SCD pts. received BT for respiratory deterioration in ICU. | n/a                  | 0 (0)                | n/a                    | n/a                    |
| Oudhia et al. [101] | USA     | 4 (n/a)           | n/a         | n/a             | None          | None          | SCD pts. had a high risk for severe disease course and high case-fatality rate in comparison to the US population, and non-SCD pts. of the same age showed lower mortality. | n/a                  | 0 (0)                | n/a                    | n/a                    |
| Panepinto et al. [30] | France | 128 (77/51)       | 29          | HbSS or HbSβ+ (135), HbSC or HbSβ+ (42), Unknown (1) | BT (68) | None          | SCD pts. had relatively mild disease course. Black SCD pts. are more likely to be hospitalized and develop pneumonia and pain than Black pts. w/o SCD but do not differ in mortality. | 128 (68.5)            | 13 (7.3)              | n/a                    | n/a                    |
| Ramachandran et al. [31] | USA     | 5 (5/4)           | 20          | HbSS (6), HbSC (1)  | HU (6), BT (6) | COVID-19 w/o SCD | SCD pts. had relatively mild disease course. Black SCD pts. are more likely to be hospitalized and develop pneumonia and pain than Black pts. w/o SCD but do not differ in mortality. | n/a                  | 0 (0)                | 3 (5.6)                | 10 (3.2)               |
| Singh et al. [33] | USA     | 312 (117/195)     | 31          | HbSS or HbSβ+ (129), HbSC, HbSβ+ or HbSE (37) | Matched COVID-19 pts. w/o SCD | None          | Significant number of hemoglobinopathy pts. in the UK developed COVID-19; SCD pts. in the age groups 18–49 and 50–79 have an increased risk of COVID-19 related death. Pregnant women with SCD have increased risk of pregnancy complications in comparison to non-SCD pregnant women. | 129 (41.3)           | 10 (3.2)              | 60 (19.2)              | 19 (6.6)               |
| Telfer et al. [32] | UK      | 166 (71/95)       | n/a         | HbSS or HbSβ+ (6), HbSβ+ (1) | BT (60) | None          | SCD pts. had relatively mild disease course. Black SCD pts. are more likely to be hospitalized and develop pneumonia and pain than Black pts. w/o SCD but do not differ in mortality. | 128 (77)             | 11 (6.6)              | n/a                    | n/a                    |
| Waymire et al. [94] | India   | 7 (0/7)           | 28          | HbSS (6), HbSβ+ (1) | None          | Pregnancy COVID-19 w/o SCD | SCD pts. had relatively mild disease course. Black SCD pts. are more likely to be hospitalized and develop pneumonia and pain than Black pts. w/o SCD but do not differ in mortality. | n/a                  | 0 (0)                | 11 (0.7)               | 11 (0.7)               |

Abbreviations: ACS, acute chest syndrome; AKI, acute kidney injury; ALL, acute liver injury; BT, blood transfusion; CKD, chronic kidney disease; DBP, diastolic blood pressure; ED, emergency department; h/o, history of; Hosp., hospitalization; HTN, hypertension; HU, hydroxyurea; ICU, intensive care unit; IMV, invasive mechanical ventilation; LOS, length of stay; n/a, not available, not applicable, unknown, or not specified; NIV, non-invasive ventilation; pts., patients; SCD, sickle cell disease; SCT, sickle cell trait; VOC, vaso-occlusive crisis; w/o, without.

1 Focus of study was not SCD.
2 50 total patients with a total of 67 VOC episodes, hospitalization rate is based on total episodes.
3 In-hospital.
4 Pediatric arm of the study.
5 Adult arm of the study.
### Table 3

Case reports of patients with SCD and COVID-19. Authors listed in alphabetical order.

| Authors                  | Country       | Sample size | Sample age (years) | Hemoglobin genotype | SCD therapy (n) | Main findings                                                                                                                                                                                                 |
|--------------------------|---------------|-------------|-------------------|--------------------|-----------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Abdulfaroukh et al.      | Bahrain       | 6 (4/2)     | 31                | HbSS (6)           | HU (6)          | Infection rate, clinical course, and viral clearance of SCD pts. were no different from pts. w/o SCD.                                                                                                            |
| Al-Hebshi et al.         | Saudi Arabia  | 2 (1/1)     | 13                | HbSS (2)           | HU (2) BT (1)   | VOC and ACS complicated infection but were effectively treated according to national guideline and standard practice.                                                                                       |
| Allison et al.           | USA           | 1 (1/0)     | 27                | HbSC (1)           | BT (1)          | Early red blood cell exchange treatment may have helped mitigate COVID-19 pneumonia and ACS.                                                                                                                  |
| Al-Naami et al.          | Kuwait        | 3 (2/1)     | 25                | HbSS (3)           | HU (1)          | Disease course was mild to moderate w/o complications or death.                                                                                                                                              |
| Al-Naami et al.          | Saudi Arabia  | 1 (1/0)     | 29                | HbSS/Siβ+         | HU (1) BT (2)   | Pt presented asymptomatic with h/o sore throat and loose motions; Uneventful recovery.                                                                                                                     |
| Al Sabahi et al.         | Oman          | 5 (4/1)     | 5                 | HbSS/HbSβ+        | HU (2) BT (4)   | Variable disease courses. All cases recovered.                                                                                                                                                              |
| Al Yazidi et al.         | Oman          | 7 (n/a)     | n/a               | n/a                | n/a             | Most pts. had SCD-related symptoms, including ACS, VOC, and splenic sequestration. Three required supplemental oxygen. ICU admission for rapid respiratory degradation after initial phase with mild symptoms. |
| André et al.             | France        | 1 (0/1)     | 5                 | n/a                | n/a             | ICU admission for rapid respiratory degradation after initial phase with mild symptoms.                                                                                                                     |
| Asshim et al.            | USA           | 11 (4/7)    | 44                | HbSS (5), HbSC (4), HbSβ+ (1), HbSβ+ (1) | BT (1)          | Pt presentations varied from mild to severe; Older age/milder genotypes had worse outcomes; ACS and COVID-19 pneumonia present similarly and should be differentiated before treatment. |
| Appiah-Kubi et al.       | USA           | 7 (2/5)     | 14                | HbSS (6), HbSC (1) | HU (4) BT (1)   | One ICU stay, no intubations. Favorable outcomes were due to early diagnosis, use of antivirals, anti-inflammatory agents, and anticoagulants.                                                                    |
| Argüello-Marina et al.   | Spain         | 7 (n/a)     | n/a               | n/a                | n/a             | Two ICU admissions, one patient required ventilation. Variable medications and treatments used. Overall, patients had short lengths of stay and no deaths.                                                           |
| Azared et al.            | Belgium       | 3 (1/2)     | 30                | HbSS (3)           | HU (3) BT (3)   | SCD pts. showed mild clinical presentation of COVID-19, which can be misleading; If BT is performed, always consider risk of DHTR.                                                                          |
| Beerkens et al.          | USA           | 1 (1/0)     | 21                | HbSβ (1)          | HU (2) BT (1)   | COVID-19 may have caused or exacerbated ACS.                                                                                                                                                                |
| Chakravorty et al.       | UK            | 10 (3/7)    | 37                | HbSS (10/10)      | HU (2)          | All but one seemed to be experiencing a relatively mild course despite significant comorbidities.                                                                                                |
| Chen-Goodspeed et al.    | USA           | 5 (3/2)     | 31                | HbSS (4), HbSC (1) | HU (3) BT (2)   | SCD pts. did not present typical COVID-19 symptoms but rather more typical SCD symptoms, including VOC; SCD pts. presenting with VOC should immediately be tested for COVID-19. |
| Dagalakas et al.         | USA           | 1 (1/0)     | 0.5               | HbSS              | n/a             | Pediatric pt. presented with mild symptoms and remained stable throughout hospitalization.                                                                                                                                 |
| De Luna et al.           | France        | 1 (1/0)     | 45                | HbSS              | BT (1)          | Severe pneumonia and severe ACS effectively treated with hydroxychloroquine and tocolzumah.                                                                                                                                 |
| Elia et al.              | Brazil        | 3 (1/2)     | 11                | HbSS (2), HbSC (1) | HU (2)          | SCD pts. showed better clinical evolution than predicted; Pulmonary viral infections can trigger SCD symptoms and need for hospitalization; COVID-19 symptoms are very similar to ACS symptoms, which affects clinical decision. |
| Ehrbler et al.           | USA           | 1 (0/1)     | 39                | HbSS              | BT (1)          | Acute crisis and COVID-19 pneumonia effectively treated with vasoconstr in lieu of single BT.                                                                                                                      |
| Espanol et al.           | USA           | 1 (0/1)     | 8                 | HbSS              | HU (1) BT (1)   | Patient was successfully managed after developing severe ACS secondary to COVID-19, complicated by cortical vein thrombosis and multisystem inflammatory syndrome.                                           |
| Fronza et al.            | Italy         | 1 (0/1)     | 44                | n/a               | BT (1)          | Patient recovered and was discharged after severe anemia and acute lung failure.                                                                                                                                 |
| Hardy et al.             | Ghana         | 3 (0/3)     | 28                | HbSS (1), HbSC (2) | BT (2)          | One had severe COVID-19 pneumonia. Two had severe SCD crisis. All had favorable outcomes.                                                                                                                     |
| Heilbronner et al.       | France        | 4 (1/3)     | 15                | HbSS (4)          | HU (2) BT (4)   | All pts. presented with ACS and admitted to PICU. Pts were effectively treated with erythropoiesis, NIV, and typical supportive treatment.                                                                     |
| Hussain et al.           | USA           | 4 (2/2)     | 33                | HbSS (2), HbSC (1), HbSβ+ (1) | HU (3) BT (1)   | All four pts. initially presented to the ED for typical VOC; the clinical course of their COVID-19 infection was rather mild.                                                                                |
| Jacob et al.             | USA           | 1 (1/0)     | 2                 | HbSS              | HU (1) BT (1)   | Pediatric pt. with severe anemia, splenic sequestration crisis and COVID-19; Pt greatly improved following BT. Mild clinical course during hospitalization, with no apparent morbidity related to COVID-19. |
| Justino et al.           | Brazil        | 1 (0/1)     | 35                | HbSS              | BT (1)          | Despite previous h/o pulmonary disease and current pregnancy, clinical course was very favorable.                                                                                                          |
| Kasinathan et al.        | USA           | 1 (0/1)     | 20                | HbSC              | BT (1)          | SCD diagnosis was unknown at time of presentation. Pulmonary embolism complicated condition.                                                                                                               |
| Martone et al.           | USA           | 1 (0/1)     | 19                | HbSβ+             | BT (1)          | PT developed ACS with fever, decreased hemoglobin, but no hypoxemia, which was managed with BT and mortality was 0.                                                                                     |

(continued on next page)
addition to perhaps COVID-19 triggering an acute vaso-occlusive crisis requiring hospitalization, as well as overlap in SCD and COVID-19 among 587,554 patients without SCD, which represents an overall mortality rate of 4.6%. In the same cohort studies, there were 21,986 COVID-19-related fatalities among 587,554 patients without SCD, which represents an overall mortality rate of 3.7%.

### 3.4. Risk of death

Among all observational cohort studies with adult patients included in this review, there were 79 COVID-19-related fatalities among 1720 patients with SCD, which represents an overall mortality rate of 4.6%. In the same cohort studies, there were 21,986 COVID-19-related fatalities among 587,554 patients without SCD, which represents an overall mortality rate of 3.7%. Based on this data from 7 different countries (i.e., Bahrain, France, India, Oman, Turkey, UK, and USA) SCD, relative to non-SCD, is associated with an overall 1.2-fold increased risk of COVID-19-related death among adult patients. In line with this observation, one of the largest cohort studies with SCD patients to date [38], conducted in the UK with a non-SCD comparison group and covariate-adjusted analysis, showed that patients with SCD who were 2.5-fold more likely to die from COVID-19 than patients without SCD at any time during the study (hazard ratio (HR) = 2.55, 95% CI = 1.36 to 4.75). Hazard ratio not only considers the total number of events, such as in relative risk and odds ratio estimates, but also the timing of each event [51]. Interestingly, Hippisley-Cox et al. [52] analyzed vaccinated cohorts and reported a substantially higher hazard ratio (HR = 7.73) for SCD patients than Clift patients.
et al. 2021 [38] (HR = 2.55) who analyzed mostly unvaccinated people in the same UK database. This may suggest that persons with SCD have a suboptimal response to COVID-19 vaccination, which warrants further investigation.

There are additional cohort studies that reported a higher risk of COVID-19 related death associated with SCD [30,32,36,37], although these studies lacked proper comparison groups. It is possible that negative outcomes reported in some of these studies are associated with other factors, such as access to care, pre-existing conditions, and other risk factors unique to these study samples, rather than SCD itself [36]. Among case studies, nearly all studies reported favorable outcomes, regardless of disease course. In 3 case studies, there were 4 deaths (all adults) out of 121 individuals with SCD [42,53,43], which represents 3.3% mortality rate.

Interestingly, SCD cohort studies utilizing controls matched for pre-existing conditions reported no significant differences in COVID-19-related mortality rate between patients with or without SCD [39,52], or between SCD patients with or without COVID-19 [26]. This implies that SCD patients do not have different COVID-19-related mortality rate as compared to non-SCD patients who have similar rates of comorbidity and end-organ damage. It further indicates that organ damage, caused by SCD or another condition, is a risk factor for COVID-19-related death.

Even though SCD is associated with increased risk of COVID-19 related death, it may not be as high as feared. While viral infections can trigger SCD symptoms [50,54], favorable COVID-19 outcomes for patients with SCD may in part be explained by the high hospitalization rate for this population, which could have contributed to timely intervention for symptomatic patients and improved survival probability. It is also possible that certain pathophysiological characteristics of SCD provide protective effects from fatal COVID-19 disease [55]. In line with this idea, some case studies reported that those with milder hemoglobin genotype had worse outcomes [42,56], although this could also be explained by a lack of hydroxyurea treatment associated with milder genotypes, which may have contributed to unfavorable outcomes. It has been recently demonstrated by proteomic analysis that neutrophils in patients with SCD present an unexpected activation of the interferon-α signaling pathway [57]. Interferons have been suggested as being a potential treatment for COVID-19 due to its antiviral activities [58,59]. The notion that SCD has some protective effect against COVID-19 should be further explored.

We conclude that based on the available empirical data in the literature, adults with SCD have an increased risk of death and a different temporal progression of death or survivorship curves following COVID-19 diagnosis as compared to adults without SCD [39,52], or between SCD patients with or without COVID-19 [26]. This implies that SCD patients do not have different COVID-19-related mortality rate as compared to non-SCD patients who have similar rates of comorbidity and end-organ damage. The effects of demographic, clinical, and socioeconomic variables on risk estimates of critical illness and death are further discussed in section 3.5 Risk factors. Of important note, although some studies have reported favorable short-term outcomes among COVID-19 patients with SCD, the long-term effects of SARS-CoV-2 infection are unknown. Patients with COVID-19 and SCD may experience lingering symptoms and long-term health problems that differ from the general population. Therefore, long-term follow-up studies of these individuals are warranted. Furthermore, there is limited data on less severe and other types of COVID-19-related outcomes among SCD patients. One cohort study reported that persons with SCD experience more COVID-19-related pneumonia and pain than individuals without SCD [53].

3.5. Risk factors

Observational cohort studies [27,29,32,36] and case reports [43,53] suggest that advanced age, pre-existing conditions, and male sex are risk factors for unfavorable COVID-19 outcomes in patients with SCD, similar to the reported risk factors in the general population [60,61]. Furthermore, because of marked differences in disease severity between different sickle genotypes, COVID-19 outcomes may vary between sickle cell patients. Interestingly, various studies observed that genotypes associated with milder SCD (i.e., HbSC, HbSE, HbSβ+) had no different or worse outcomes than genotypes associated with more severe SCD (i.e., HbSS, HbSβ0) [27,30,32,42,56]. We speculate that sickle cell anemia-specific therapies or pathophysiology, such as activation of the interferon-α signaling pathway as discussed earlier in this work, may provide a protective effect against COVID-19, but the reasons for this observation are unknown and require further investigation. It is known, however, that SCD symptoms could complicate the COVID-19 disease course. For example, Mucalo et al. [29] reported that adults with a history of pain are at higher risk of worse disease severity. Another major concern unique to patients with SCD is pulmonary thrombosis, which is prevalent in both SCD [62] and COVID-19 [63]. Therefore, having both conditions could conceivably result in even higher risk of severe disease. Persons with SCD would benefit from individual risk assessment for poor COVID-19 outcomes although a history of SCD complications does not necessarily lead to unfavorable outcomes. For example, Ansum et al. [42] reported that prior sickle cell complications such as avascular necrosis of the joints, hypersplenism requiring splenectomy, and cerebrovascular accident did not associated to the outcome of patients with COVID-19.

3.6. SCD-modifying therapy

Several authors have speculated on the potential beneficial effects of SCD-directed therapies against COVID-19 disease. Simple or exchange blood transfusion was the primary therapy reported by observational cohort studies [26-32,35-37,40,64-66]. Among case studies, 47% of SCD patients (60 out of 129 individuals) received regularly scheduled chronic transfusion prior to COVID-19 diagnosis and/or received transfusion during COVID-19 disease course. Several reports suggest that early red blood cell exchange is successful in the treatment of severe COVID-19 pneumonia [35,54,55,67], severe hemolysis and vaso-occlusive crisis [68], and might prevent patients with SCD from experiencing further clinical deterioration [69] and intubation [67].

Hydroxyurea treatment is another common therapy for SCD [27,29,31,35-37,40,70] and was received by 32% of SCD patients (41 out of 129 individuals) in case studies. The beneficial effects of hydroxyurea treatment on SCD-related morbidity and mortality are well known [71,72]. Minniti et al. [36] reported that all deaths occurred in patients not on hydroxyurea or other SCD-modifying therapies. Morrone et al. [73] reported that patients on hydroxyurea did not develop ACS. Mucalo et al. [29] though found that hydroxyurea had no effect on hospitalization or COVID-19 disease severity. Interestingly, of the 4 reported case-study deaths, 3 did not receive transfusion or hydroxyurea treatment, and one patient did not have data on transfusion or hydroxyurea treatment. Prospective studies are needed to further examine the effects of SCD-modifying therapies on COVID-19 outcomes.

3.7. Pediatric patients with SCD

In the general population, children are less likely to experience severe COVID-19 outcomes as compared to adults [74], but only few studies [29,70] have examined children with SCD and SARS-CoV-2 infection in sufficiently large cohorts from which valid inferences can be drawn, as the vast majority of studies were case reports with less than 10 patients [56,65,66,73,75-89].

In one of the largest pediatric studies to date, Mucalo et al. [29] analyzed 364 US children with SCD and SARS-CoV-2 infection (median age 11 years, 72% HbSS/HbS-beta thalassemia, 56% on hydroxyurea treatment) and found most children to be asymptomatic (25.5%) or to experience mild to moderate symptoms (65.6%) although some had more severe or critical symptoms (8.2%). Furthermore, the sample was associated with 40% hospitalization rate, 5.8% ICU admission, 1.1% ventilator use, and 1 death (0.3%). Risk factors for worse COVID-19
outcomes were history of pain, and renal and heart/lung comorbidities. No effects of hydroxyurea on hospitalization or COVID-19 severity were observed. These findings were mostly in accordance with a smaller French cohort study by Broussy and colleagues [70] who analyzed 39 children with SCD and SARS-CoV-2 infection (median age 12 years, 90% HB-SS, 64% on hydroxyurea treatment) and found that none had displayed symptoms except one who was hospitalized for mild vaso-occlusive symptoms with a favorable outcome. It was hypothesized that activation of the type I Interferon (IFN-I) pathway may have contributed to partial protection from SARS-CoV-2 severe illness and death seen in their pediatric sample. Among the smaller pediatric studies [56,65,66,73,75-89] that reported on a combined 42 children with SCD, there was 1 (2.4%) patient with a history of prior pulmonary and cardiac complications who died from COVID-19 disease.

While the literature suggests that most children with SCD who become infected with SARS-CoV-2 have mild disease and low risk of death, children with SCD, especially those with SCD-related comorbidities [29,65,66,90], are more likely to be hospitalized and require escalated care than children without SCD of the same age [91]. However, compared to adults with SCD [29] and adults without SCD [38], children with SCD are less likely to experience COVID-19-related severe illness and death. There was not sufficient data to review children with SCT and COVID-19. Multicenter studies with larger sample size are needed to confirm these preliminary findings.

4. Patients with COVID-19 and sickle cell trait

Table 4 summarizes the study parameters and main findings of 7 observational cohort studies [28,33,38,39,92-94] and 4 case studies [75,90,95,96] reporting on a total of 1937 individuals with SCT and COVID-19.

4.1. Sample characteristics

There were 1937 persons with SCT and SARS-CoV-2 infection analyzed by all studies combined and included in this review (Table 1). The SCT cohort had a median age of 54 years (interquartile range = 40 to 61 years) and was 54% female. A relatively large proportion of patients had hypertension (42%), followed by asthma/COPD (30%) and diabetes (23%), which may represent an age-related effect. Patients presented with 96% oxygen saturation, 100.3F temperature, and a BMI of 32. All laboratory values where within normal range, except for lactate dehydrogenase, c-reactive protein, and D-dimer which were above normal range possibly due to systemic inflammation and coagulation dysfunction associated with more severe COVID-19 [97,98]. Of note, aggregated data, including laboratory values, should be noted, aggregated data, including laboratory values, should be

Table 4

Study characteristics and main findings of reports on individuals with SCT and COVID-19. Authors listed in alphabetical order.

| Authors | Country | Study type | Sample size (M/F) | Age (years) | Control group | Main findings | SCT | Non-SCT |
|---------|---------|------------|------------------|-------------|---------------|---------------|-----|---------|
| Al-Hebshi et al. [73] | Saudi Arabia | Case report | 1 (0/1) | 50 | None | No significant symptoms, except for headache and fatigue prior to testing. Normal labs and chest X-ray. | 0 (0) | 0 (0) | n/a |
| Balanchivadze et al. [28] | USA | Case report | 18 (3/15) | 58 | None | SCT pts. generally had mild disease course with lower chances of intubation, ICU admission, and death. | 11 (61) | 1 (6) | n/a |
| Clift et al. [38] | UK | Obs. | 1346 (n/a) | n/a | COVID-19 w/o SCT | SCT pts. had 1.4-fold and 1.5-fold increased risk of hospitalization and death due to COVID-19, respectively. | 98 (7.3) | 50 (3.7) | 23,561 (4.4) | 19,008 (3.5) |
| Hoogenboom et al. [39] | USA | Obs. | 62 (13/49) | 47 | Matched and unmatched COVID-19 w/o SCT | SCT pts. did not differ from (un)matched controls in laboratory values or outcomes. | 31 (50) | 7 (23) | n/a | 20 (18) |
| Merz et al. [93] | USA | Obs. | 20 (11/9) | 66 | COVID-19 w/o SCT | SCT did not impact respiratory, renal, or circulatory complications or mortality in COVID-19 pts. when compared to non-SCT pts. | n/a | 3 (15) | n/a | 19 (13) |
| Quaresima et al. [87] | Italy | Obs. | 2 (1/1) | 51 | None | Prolonged positivity for SARS-CoV-2, but mostly asymptomatic to mild disease course. | 0 (0) | 0 (0) | n/a |
| Resurreccion et al. [92] | UK | Case report | 14 (6/8) | 64 | COVID-19 w/o SCT | Black SCT pts. had similar infection rates, but higher mortality compared to Black pts. w/o SCT. Diabetes was a risk factor for COVID-19 related death. | n/a | 4 (29) | n/a | 21 (13) |
| Sheha et al. [95] | Egypt |Obs. | 1 (0/1) | 22 | None | First case of SCD diagnosed due to concurrent COVID-19. | n/a | 0 (0) | n/a |
| Singh et al. [123] | USA | Case report | 449 (237/212) | 37.7 | Matched COVID-19 pts. w/o SCT | Black pts. with SCT did not differ in COVID-19 disease course or outcomes compared to Black pts. w/o SCT. | 79 (18) | 10 (2.2) | n/a |
| Tafti et al. [96] | USA | Obs. | 1 (1/0) | 33 | None | Pt developed rhabdomyolysis, myonecrosis, and an abscess, all of which were exacerbated by COVID-19. | n/a | 0 (0) | n/a |
| Waghmare et al. [124] | India | Obs. | 24 (0/24) | n/a | Pregnant COVID-19 w/o SCT | Pregnant women with SCT have increased risk of pregnancy complications in comparison to pregnant women w/o SCT. | n/a | 1 (4.2) | n/a | 11 (0.7) |

Abbreviations: ACS, acute chest syndrome; AKI, acute kidney injury; ALI, acute liver injury; ARDS, acute respiratory distress syndrome; CT, computed tomography; ED, emergency department; Hosp., hospitalization; ICU, intensive care unit; IMV, invasive mechanical ventilation; LOS, length of stay; n/a, not available, not applicable, unknown, or not specified; Mort., mortality; pts., patients; Obs., observational cohort study; SCT, sickle cell disease; COVID-19, sickle cell trait; w/o, without.

1. In-hospital.
2. Patient tested positive for SCT but was suspected SCD.
interpreted with caution due to small sample sizes.

4.2. Clinical course

Most studies \([28,39,33,75,87,93]\) reported mild COVID-19 disease course for individuals with SCT similar to individuals without SCT, and only very few SCT patients required escalated care (i.e., intensive care, intubation, need for mechanical ventilation) or developed severe disease complications. However, pregnant women with SCT may experience increased risk of pregnancy complications (e.g., gestational hypertension, intrauterine growth restriction) and more COVID-19-related symptoms in comparison to pregnant women without SCT \([94]\). Among case studies, only 2 patients with SCT reportedly had more severe disease course. In one case, the patient was not known to have SCT prior to hospitalization for COVID-19 and may have had disease phenotype masked by the transfusions \([95]\). The second SCT case had rhabdomyolysis after significant exertion due to exercise \([96]\). Some studies did not collect or analyze any clinical course data from its cohort \([38,92]\).

4.3. Hospitalisation trends

Hospitalization rates for patients with SCT and COVID-19 varied greatly between studies from 17.6% \([33]\) to 61.1% \([28]\), while 1 study did not report it for their cohort \([92]\). In a large UK cohort, SCT was associated with a 1.4-fold increased risk of hospitalization compared to the general population adjusted for age, sex, and ethnicity \([38]\). Variation in reported hospitalization rates is likely due to differences in cohort characteristics as studies with higher hospitalization rates included significantly older patients with SCT who had many comorbidities \([28,39]\), which can be viewed as additional risk factors for hospitalization. In fact, studies with comorbidity-matched controls showed no differences in COVID-19-related hospitalization for those with SCT compared to those without SCT \([39,33]\), which implies that comorbidities and not SCT per se are associated with an increased risk of admissions.

4.4. Mortality rate

Similar to hospitalization rates, the reported COVID-19-related mortality rates for persons with SCT vary greatly between studies from 2.2% \([33]\) to 28.6% \([92]\). In one of the largest studies with SCT patients to date, persons with SCT were found to have a 1.5-fold increased risk of death due to COVID-19, similar to SCD albeit to a lesser extend \([38]\). Another study confirmed these findings \([92]\), but only in a small sample without demographic-matched controls. There are also large cohort studies that have shown no difference in mortality rates between individuals with or without SCT and COVID-19 \([39,33,93]\). Differences in matching criteria, especially related to comorbidity-matching, may explain some of the discrepancies observed in reported mortality rates between studies. No studies analyzed risk factors for mortality except for one \([92]\), which found preexisting diabetes to increase risk of death in SCT patients. Unique factors that lead to severe disease and death among SCT patients, such as comorbidity burden or genetic predisposition, are unknown and require further investigation.

5. Summary and conclusions

This comprehensive review summarizes the clinical characteristics and COVID-19-related outcomes of patients with SCD and SCT from the published literature to date. While most studies reported mild to moderate COVID-19-related disease course in this patient population, the literature suggests that SCT is associated with an increased risk of hospitalization and death from COVID-19, unless compared to controls with similar comorbidities and end-organ damage. Advanced age, preexisting conditions, and male sex are risk factors for unfavorable COVID-19 outcomes in patients with SCD, similar to the reported risk factors in the general population. There is some evidence that SCD-modifying therapies such as transfusion and hydroxyurea may be associated with better COVID-19 outcomes, but prospective studies are needed for confirmation. There is some evidence that persons with SCT have increased risk of COVID-19-related hospitalization and death although more studies with risk-stratification and properly matched controls are needed to confirm these findings.

6. Future perspectives

Based on this review of the current literature, there are various suggested future directions to move the field forward and improve our understanding of COVID-19 outcomes among patients with SCD and SCT. First, an important focus area for future studies should include multi-center collaboration to generate larger cohorts, which will generate more definite evidence related to COVID-19 outcomes among patients with SCD/SCT. Currently, two-thirds of published COVID-19 studies with SCD/SCT patients are case reports. While case reports provide important and detailed information at the patient level, often detecting novelties and generating hypotheses, the lack of generalizability, danger of over-interpretation, and publication bias are major limitations of the case report genre \([99]\). For this reason, population level inferences should be based on larger controlled experimental studies. Second, into what extend hemoglobin genotype or sickle cell therapy impact COVID-19 infection is largely unknown. Because of marked differences in disease severity between different hemoglobin genotypes, COVID-19 outcomes may vary between sickle cell patients, which requires further investigation. Third, future studies should include case-matched controls that account for the unique sample characteristics of SCD and SCT populations. Patients with SCD and SCT often have unique underlying conditions that could make them more vulnerable to poor COVID-19 outcomes. Fourth, while most studies have reported favorable short-term outcomes among COVID-19 patients with SCD and SCT, especially related to severe illness and death, the long-term effects of SARS-CoV-2 infection are unknown. It is possible that SCD patients with prior SARS-CoV-2 infection experience more lingering symptoms and long-term health problems compared to survivors of COVID-19 disease in the general population. Therefore, long term follow-up studies with longitudinal assessment of post-COVID-19 symptoms are warranted.

Practice points

- Patients with SCD infected with SARS-CoV-2 should be followed very closely as they often have underlying conditions that could make them more vulnerable to poor COVID-19 outcomes.
- A low threshold for admission of SCD patients is common as SARS-CoV-2 infection could trigger acute vaso-occlusive crisis requiring hospitalization and due to overlap of SCD and COVID-19 symptoms (e.g., the clinical picture for both ACS and COVID-19 pneumonia can be similar).
- Transfusion and hydroxyurea treatment have demonstrated benefit in treating SCD-related morbidity and mortality. The role of SCD-modified therapies in COVID-19 appears to be associated with favorable outcomes and is under active investigation.
- While most SCD patients only experience mild to moderate COVID-19 symptoms, similar to the general population, SCD is associated with an increased risk of COVID-19-related hospitalization and death, which indicates that serious illness and death still occurs especially among vulnerable individuals in this patient group.
- Most COVID-19 patients with SCT appear not different from the general population with COVID-19 in terms of admission laboratory values, clinical course, hospitalization trends, and mortality, but additional studies are needed to confirm these findings.
Research agenda

- Larger, multisite, randomized studies are needed to generate more definitive evidence related to COVID-19 outcomes among patients with SCD/ SCT.
- Because of marked differences in disease severity between different hemoglobin genotypes, COVID-19 outcomes may vary between sickle cell patients, which requires further investigation.
- Inclusion of properly matched controls is important to account for the unique sample characteristics of SCD and SCT populations.
- Long term follow-up studies of patients with SCD/SCT are needed as the long-term effects of COVID-19 may affect these patients differently as compared to the general population.

Author contributions

All authors (WSH, TTA, DMM, NB, GD, WBM, DM, and TD) made substantial contributions to all of the following: (1) the conception and design of the study, acquisition of the data, and analysis and interpretation of the data, (2) drafting the article or revising it critically for important intellectual content, (3) review and approval of the final manuscript submitted.

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Declaration of competing interest

The authors report no competing interests.

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