A pilot trial of human amniotic fluid for the treatment of COVID-19

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Abstract

Objective: Vertical transmission from SARS CoV-2-infected women is uncommon and coronavirus has not been detected in amniotic fluid. Human amniotic products have a broad immune-mediating profile. Observing that many COVID-19 patients have a profound inflammatory response to the virus, we sought to determine the influence of human amniotic fluid (hAF) on hospitalized patients with COVID-19.

Results: A 10-patient case series was IRB-approved to study the impact of hAF on hospitalized patients with documented COVID-19. Nine of the 10 patients survived to discharge, with one patient succumbing to the disease when enrolled on maximal ventilatory support and severe hypoxia. The study design was altered by the IRB such that the last 6 patients received higher dose of intravenous hAF. In this latter group, patients that had observed reductions in C-reactive protein were associated with improved clinical outcomes. No hAF-related adverse events were noted. Acknowledging some of the inherent limitations of this case series, these results inform and catalyze a larger scaled randomized prospective trial to further investigate hAF as a therapy for COVID-19.

Trial Registration ClinicalTrials.gov: NCT04319731; March 23, 2020

Keywords: COVID-19, Amniotic fluid, Inflammation, C-reactive protein

Introduction

Early data in the COVID-19 pandemic suggested that vertical transmission from SARS CoV-2-infected women is uncommon and virus has not been detected in the amniotic fluid [1–4]. These observations raise the question of whether or not there is something intrinsically protective in the fluid. Indeed, human amniotic products have a broad immune mediating profile [5]. Human amniotic membrane (hAM) and human amniotic fluid (hAF) have been shown to reduce inflammation, have antimicrobial properties, and confer a low risk of immunogenicity [6–11]. Purified hAF is non-antigenic solution devoid of any cellular products (i.e., it is not to be confused with umbilical cord-derived, AF-derived stem cell products, or AF embolism) that nature developed [5, 12]. Hence, amniotic products make ideal biocompatible scaffolds for the treatment of diverse conditions, including but not limited to intractable epithelial defects, burns, diabetic/vascular ulcers, partial limbal cell deficiencies, peripheral nerve regeneration, tendon repair, and Stevens-Johnson syndrome [13–15].

Our group has developed experience with the clinical use of amniotic products and is currently using hAM as a tissue allograft in burn patients, for digital ulcers in scleroderma patients, as a nerve wrap to protect nerves from adhesions post-surgery, and as a trachea-stenosis-tracheal stent covering. Additionally, our burn unit has successfully injected purified hAF into more than 500 burn patient wounds to augment graft survival. hAF is also currently being used experimentally under IND approval to treat ocular graft versus host disease and ocular PRK.
Given the use of amniotic products for the treatment of inflammation in other conditions, our group proposed an experimental and innovative use of hAF as its impact on cardiopulmonary failure has not been previously investigated. In light of the profound local and systemic inflammation associated with COVID-19 [16], it is our global hypothesis that hAF could significantly mitigate the progression of disease. An expedited protocol was approved by the University of Utah Institutional Review Board to study the influence of hAF in 10 patients with confirmed COVID-19. Herein, we report the results of this case series.

Main text

Purified hAF is a cell-free, non-antigenic solution (not to be confused with umbilical cord-derived or AF-derived stem cell products) that is processed and manufactured at the University of Utah for clinical use [8, 12]. The initial trial design identified two cohorts of hospitalized, symptomatic, and laboratory verified SARS CoV-2 patients: (1) mechanically ventilated patients were administered hAF both intravenously (3 cc) and nebulized (3 cc) for 5 days; and (2) non-mechanically ventilated patients were administered only nebulized (3 cc) hAF. After the first 4 patients, concerns with utilizing aerosolized therapies in COVID-19 patients required a temporary hold and subsequent re-design. Thereafter (the last 6 patients), all received a higher dose of intravenous hAF (10 cc) for 5 consecutive days. Patients were eligible for this study if there were over the age of 18, had a SARS CoV-2 laboratory positive test obtained within 14 days of enrollment, were hospitalized, and symptomatic (e.g., cough, fevers, shortness of breath, sputum production). There were no exclusion criteria for the case series. Furthermore, to the best of our knowledge, there are no absolute contraindication for the clinical use of our purified hAF. Participants were enrolled over the span of seven weeks between late March and early May of 2020.

The demographics (Table 1) of the total 10 patient cohort included the following: 40% female; average age of 51.9 years old (range 24–76); five White, two American Indian/Alaskan Native, two Hispanic, and one as unknown/other. Average weight was 91 kg (range 42–140). Eight of the 10 participants had underlying comorbid conditions, of which the most common were diabetes (n = 6) and hypertension (n = 5). As this is a small case series, pertinent information is provided in a patient-by-patient manner (Table 1). Of the 10 patients enrolled, seven were admitted directly to the intensive care unit (ICU). All ten patients required oxygen support during their hospital admission. Three patients required a nasal cannula or simple mask as their highest level of support, while two were placed on a high flow nasal cannula as their highest support. Five patients required mechanical ventilation. One patient (#1) was enrolled on high-level ventilatory support (100% FiO2, 18 mmHg PEEP) in the prone position and ultimately required 41 days of veno-veno extracorporeal membrane oxygenation (initiated after 5th day of hAF treatment) as well as tracheostomy, which was removed prior to discharge. Another patient (#6) required a tracheostomy and was discharged with this still in place, although he was no longer on mechanical ventilation. Of the nine patients who have been discharged alive, 7 required home oxygen use.

Results of primary outcome measures included hospital length of stay, ICU length of stay, ventilator-free days, and supplemental oxygen needs at the time of discharge (Table 2). The 4 patients who were mechanically ventilated and ultimately discharged alive averaged 13.5 ventilator-free days (range 5–28). With regard to supplemental oxygen use at the time of discharge, 2 patients were entirely on room air, 6 were requiring continuous oxygen, and 1 required intermittent oxygen use. Average hospital length of stay (n = 10) was 20.4 days (range 5–56 days). Nine of 10 patients were admitted to the ICU at some point during their hospitalization. Average ICU length of stay for surviving patients (n = 8) was 17.38 days (range 2–56 days). For this subgroup of 8 patients, the average ICU-free days/length of stay on the medical floor was 5.9 days (range 1–19 days).

| Table 1 Demographics and medical history |
|------------------------------------------|
| Variable | n  |
| Age (mean, range) | 52, 24–76 |
| Female sex | 4 |
| Weight (kg, mean, range) | 91 (42–140) |
| Race/Ethnicity |  |
| White | 5 |
| American Indian/Alaskan Native | 2 |
| Hispanic | 2 |
| Unknown/other | 1 |
| Medical History |  |
| Anemia | 1 |
| Coronary artery disease | 1 |
| Diabetes | 6 |
| Elevated liver enzymes | 1 |
| Heart murmur | 1 |
| Hyperlipidemia | 2 |
| Hypertension | 5 |
| Liver disease | 1 |
| Mitral valve disease | 1 |
| Respiratory disease (e.g., asthma) | 2 |
| No past medical history | 2 |
Biomarkers were evaluated before and after treatment with hAF (Table 1, Additional file 1: Table S1). Because of the small sample size and some missing data, it is difficult to make any broad statements concerning the influence of hAF on these biomarkers. That stated, in the latter cohort that received 10 cc hAF intravenously, a mean reduction in C-reactive protein by 38% was observed. Particularly notable is an apparent decrease in markers in those patients (4/6) that had improved clinical profiles. Conversely, two of the six patients saw increases in these inflammatory biomarkers—one had a prolonged hospitalization and the other was discharged to a rehabilitation hospital. One patient (#3) died on hospital day 8. She had multiple comorbidities including morbid obesity (BMI 55 kg/m²) and was extremely ill (maximal ventilatory support) on admission, and had a noticeable increase in her biomarkers. Outside of the above-mentioned issue related to aerosolized therapy, where theoretic concerns about safety to the provider delivering the nebulized hAF were never observed, there were no reported safety concerns.

**Discussion**

The body of knowledge related to the pathophysiology and therapeutics of COVID is rapidly evolving. Our understanding of the relative contributions of innate and adaptive immune responses is similarly advancing. Indeed, many early observations have been modified as the nuances of different patient populations and presentations exist. Some patients, for example, demonstrate a syndrome consistent with cytokine storm, with a very aggressive inflammatory response. These patients might benefit from blockade of classic cytokine pathways (i.e. IL-6 antagonists). Other patients have a more subtle, perhaps less florid inflammatory response, and might not benefit as much from robust immune-modulation.

One of the attractive features of hAF is its diverse “soup” of ingredients. Expression profiling of 68 term gestation patients demonstrated that the cell-free AF transcriptome contains over 64,000 genes [5, 12]. At our center, we performed a cytokine array on AF from 17 patients with normal term pregnancy. Over 300 proteins (out of the 400 on the array) were present in AF with the majority associated with host defense and angiogenesis [12]. Biology has long taught us that blocking or augmenting a single pathway is confounded by naturally derived redundancy and alternative feedback loops. Indeed, while IL-6 is found to be upregulated in COVID patients, treatment with a singular antagonist does not necessarily change the course of the disease [17]. While some therapeutics will try to combine various mechanistic pathways[18], hAF is nature’s combination of not just one or two proteins, but rather thousands of proteins which have been mixed together and survived thousands of years.

Outside of the aforementioned studies surrounding wound healing, there is scant data with regards to the utilization of hAF for cardiopulmonary disease. Many COVID-19 patients, in addition to succumbing to acute hypoxic respiratory failure, are plagued with cardiac manifestations including myocarditis, accelerated heart failure, and arrhythmias [19]. Lung pathology specimens from the SARS-1 epidemic in 2005 demonstrated diffuse alveolar damage with extensive fibrosis [20]. Likewise, lung pathology in COVID-19 patients undergoing lung resection for lung cancer (i.e., non-autopsy pathology) revealed edema and patchy, inflammatory cellular infiltrates, particularly with macrophages [21]. Summarily, COVID-19 can cause an aggressive

| Table 2 Outcomes of 10 enrolled patients |

| Patient | Admit | Initial O2 | Maximum O2 | Co-Rx | CRP | IL-6 | D-dimer | LDH | Discharge |
|---------|-------|------------|------------|-------|-----|------|---------|-----|-----------|
| 1       | ICU   | ETT        | ECMO/Trach | HCQ   | +30%| +38%| +138%   | -27%| Alive     |
| 2       | Floor | NC         | NC         | HCQ   | N/A | N/A | N/A      | N/A | Alive     |
| 3       | Floor | NC         | ETT        | HCQ, Rem | N/A | N/A | N/A      | N/A | Deceased  |
| 4       | ICU   | NC         | NC         | HCQ   | N/A | N/A | N/A      | N/A | Alive     |
| 5       | ICU   | ETT        | ETT        | HCQ   | -84%| -60%| -21%     | -22%| Alive     |
| 6       | ICU   | ETT        | Trach      | HCQ-AZ | -91%| 0  | -42%     | N/A | Alive     |
| 7       | ICU   | HFNC       | HFNC       | HCQ   | +46%| +109%| +818%    | +109%| Alive     |
| 8       | ICU   | NC         | HFNC       | AZ    | -87%| 0  | 0        | -21%| Alive     |
| 9       | ICU   | NC         | Simple mask| HCQ   | -80%| -20%| -28%     | +109%| Alive     |
| 10      | ICU   | HFNC       | HFNC       | HCQ   | +41%| +412%| +33%     | -30%| Alive     |

Of note, protocol change (10 cc hAF I.V.) for patients 5–10. N/A not available, ECMO extracorporeal membrane oxygenation, ETT endotracheal intubation, HFNC high flow nasal cannula, ICU intensive care unit, MACE major adverse cardiac event, NC nasal cannula, O2 oxygen, Trach tracheostomy, Co-Rx Co-treatment, HCQ hydroxychloroquine, AZ azithromycin; Rem remdesivir. Biomarker are represented as a percent change between study enrollment and termination of hAF (5 days) with focus on the last 6 patients under current protocol. CRP: C-reactive protein; IL-6: interleukin-6; LDH: lactate dehydrogenase.
inflammatory and fibrotic response in both the heart and lung. This suggests that hAF, by mitigating inflammation and decreasing fibrosis, could impact the natural history of COVID-19 infected patients and provides the foundational platform for our investigation of the potential impact of systemic administration of hAF in COVID-19 patients.

Limitations
This case series has a number of limitations including the small patient cohort (thereby limiting inferential and comparative statistics), mixed patient population including those in extremis, mixed delivery protocol between the first 4 and the last 6 patient, and concomitant medical and experimental therapies (for example, hydroxychloroquine—see Table 1). In addition, this case series was performed in the initial phases of the pandemic and some of the biomarker studies in the first 4 patients were not available or collected within the timeframe set forth by protocol. Despite these significant caveats, the observations in these patients suggest that hAF can be used safely in this population and provides a potential signal of a favorable biologic influence. As such, this report provides some data to support the conceptual rationale for further investigation. This experience has propelled our application and subsequent approval from the FDA for IND status (#23,369), and has informed the design and recent implementation of a larger-scale, randomized clinical trial to evaluate the efficacy and novel utilization of human amniotic fluid as therapy for COVID-19.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s13104-021-05443-9.

Additional file 1: Table S1. Raw biomarker data before and after therapy for each individual patient: N/A: not available; CRP: C-reactive protein; IL-6: interleukin-6; LDH: lactate dehydrogenase.

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Authors’ contributions
All authors have read and approved the final manuscript. Specific roles include: Conception: CHS, JP; Design: CHS, JP, GL, NDH, IDP; Acquisition of data: CHS, CV, JP, NDH; Analysis and interpretation: All authors; Draft and revision of manuscript: CHS, JP, JET, CS, GL, NDH, IDP; All authors read and approved the final manuscript.

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Availability of data and materials
The datasets generated and/or analyzed during the current study are not publically available due to protected health information but are available from the corresponding author on reasonable request.

Ethics approval and consent to participate
Approved by the University of Utah Institutional Review Board (IRB_00131618). Consent was obtained for all enrolled (10) patients. Consent was obtained both by written and verbal means. As this trial was performed on patients with confirmed COVID-19, this approach was sanctioned by our IRB in order to decrease exposure of healthcare and research workers to the coronavirus.

Consent for publication
All patients, within the IRB-approved consent form, acknowledged and agreed to the possibility that the results of the study would be published, but specific data would be de-identified.

Competing interests
Drs. Pierce and Phillips are members of the Department of Medicine and administratively direct the Center for Translational and Regenerative Medicine (CTRM) where the human amniotic fluid is processed and subsequently delivered for clinical use. The CTRM has a number of patents with the University of Utah with regards to the utilization of human amniotic fluid for other disease processes (for example, graft versus host disease). The remaining authors have no competing interests.

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