Visceral adipose tissue in patients with COVID-19: risk stratification for severity

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Abstract

Purpose To assess visceral (VAT), subcutaneous (SAT), and total adipose tissue (TAT) estimates at abdominopelvic CT in COVID-19 patients with different severity, and analyze Body Mass Index (BMI) and CT estimates of fat content in patients requiring hospitalization.

Methods In this retrospective IRB approved HIPPA compliant study, 51 patients with SARS-CoV-2 infection with abdominopelvic CT were included. Patients were stratified based on disease severity as outpatient (no hospital admission) and patients who were hospitalized. Subset of hospitalized patient required mechanical ventilation (MV). A radiologist blinded to the clinical outcome evaluated single axial slice on CT at L3 vertebral body for VATL3, SATL3, TATL3, and VAT/TATL3. These measures along with age, gender, and BMI were compared. A clinical model that included age, sex, and BMI was compared to clinical + CT model that also included VATL3 to discriminate hospitalized patients from outpatients.

Results There were ten outpatients and 41 hospitalized patients. 11 hospitalized patients required MV. There were no significant differences in age and BMI between the hospitalized and outpatients (all \( p > 0.05 \)). There was significantly higher VATL3 and VAT/TATL3 in hospitalized patients compared to the outpatients (all \( p < 0.05 \)). Area under the curve (AUC) of the clinical + CT model was higher compared to the clinical model (AUC 0.847 versus 0.750) for identifying patients requiring hospitalization.

Conclusion Higher VATL3 was observed in COVID-19 patients that required hospitalization compared to the outpatients, and addition of VATL3 to the clinical model improved AUC in discriminating hospitalized from outpatients in this preliminary study.

Keywords Visceral adipose tissue (VAT) · CT · COVID-19

Introduction

A new RNA betacoronavirus called severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is responsible for the illness COVID-19 [1–3] that has resulted in, and continues to cause severe global morbidity and mortality. COVID-19 was classified as a global pandemic by the World Health Organization (WHO) on March 11, 2020. There are two major hypothesized mechanisms for viral infection related injury: (1) the virus binds to Angiotensin Converting Enzyme 2 (ACE2) human cell receptor which is highly expressed in type 2 alveolar cells, vascular epithelium, and cardiac myocytes; (2) a cytokine storm related to an exuberant pro-inflammatory state [4]. The spectrum of symptomatic infection ranges from mild to critical illness. Some patients with initially mild symptoms may progress over the course of a week and develop more severe disease, which may require mechanical ventilation. Pneumonia is one of the common manifestation of lung injury. Acute respiratory distress syndrome (ARDS) requiring invasive mechanical ventilation (MV) is a major pulmonary complication in patients with severe disease and can manifest shortly after the onset of dyspnea. Therefore, there is a strong need to identify and validate biomarkers that are associated with
severe symptoms and worse prognosis. Reliable prediction of high-risk patient phenotype can allow us to optimize resources and closely monitor those patients who are likely to require hospitalization, intensive care unit (ICU) admission, and MV.

Recent studies have identified a number of risk factors that predispose patients to severe illness. These include age, male sex, diabetes mellitus, hypertension, cardiopulmonary diseases, and obesity [5–7]. A number of studies have identified obesity as a risk factor not only for hospitalization but also for MV requirement in patients with COVID-19 [8]. Obesity is also associated with an increased risk of metabolic disorders such as diabetes mellitus, and cardiovascular diseases [9]. Obesity is considered a state of low-grade inflammation [10], with various inflammatory products secreted by adipose tissue. The main adipose tissue-derived inflammatory cytokines include TNFa, IL-1, and IL-6. Recent reports suggest an increased inflammatory environment leading to cytokine storm in significant number of patients with COVID-19 [11]. The mechanism of viral infection and the development of obesity seem to share some common metabolic and inflammatory reaction pathways. Infection of individuals who are already showing increased inflammation may result in higher morbidity. This could potentially provide an explanation as to why obesity is associated with worse outcome [12].

All fat depot in the body are not created equal; they confer different cardiovascular and metabolic risks [13, 14]. Subcutaneous adipose tissue (SAT) is relatively indolent, whereas visceral adipose tissue (VAT) is metabolically more active tissue that secretes inflammatory cytokines [14]. VAT is linked to metabolic syndrome, cardiovascular impairment, and increased susceptibility to infection and sepsis [15]. Hence, it is of interest to analyze the relationship between the amount of visceral fat and the severity of COVID-19 infection. VAT and SAT can be measured and classified rapidly and accurately using MRI or CT [16]. While it is possible to quantify the amount of adipose tissue within the whole abdomen, several studies have demonstrated that the single-slice measurement made rapidly at the vertebral body L3 or L4 level is an excellent estimate of the abdominal VAT and SAT in both men and women and across the spectrum of obesity [17, 18].

The aim of this study was twofold: (1) to assess VAT, SAT, and total adipose tissue (TAT) on abdominopelvic CT in SARS-CoV-2-positive patients with different severity of COVID-19 infection, and (2) to analyze body mass index (BMI) and fat content estimated at CT in patients with COVID-19 who required hospitalization.

Methods

Subjects

This study was approved by the institutional review board and was HIPAA compliant and received waiver of informed consent. A retrospective institutional database search identified all adults (≥ 18 years of age) who had an abdominopelvic CT from March 19, 2020 through April 19, 2020 with “COVID” in the CT report. This search initially identified 72 patients. Eighteen patients were excluded as COVID-19 positivity was not confirmed by the review of the electronic medical record (EMR). This resulted in the study cohort of 54 patients (38 men and 16 women; mean [± SD] age, 59.8 [14.9] years; age range 20–88 years). PCR-positive diagnosis of SARS-CoV-2 infection was confirmed in all 54 patients.

Indications for the CT were abdominal pain (n = 26), fever (n = 5), sepsis/hypotension (n = 4), acute kidney injury (n = 2), cancer (n = 2), GI bleed (n = 2), trauma (n = 3), poor oral intake/SBO/diarrhea (n = 5), vascular indications (n = 3), abnormal liver function test (n = 1), and hematuria (n = 1). Three patients were excluded due to unavailability of axial imaging for analysis (n = 1), and presence of extensive ascites (n = 2) which precludes evaluation of visceral fat (Fig. 1). The EMR was reviewed to categorize patients into different phenotypic cohorts based on disease severity: (1) patients that did not require hospital admission (outpatient), (2) patients who were hospitalized (hospitalized), and (3) subset of hospitalized patient who were admitted to the ICU and required invasive MV (hospitalized + MV). Of the 10 outpatients, 6 patients presented to ER/urgent care and 4 patients presented to outpatient imaging facility. Age, gender, and Body Mass Index (BMI) were also collected in all patients when available (Table 1). BMI was unavailable in three patients.

Some of these patients were included in two other imaging studies that focused on qualitative imaging findings at lung bases and findings of hypercoagulability in COVID-19 patients [19].

CT segmentation of VAT and SAT

De-identified axial DICOM CT images were parsed and loaded to the NIH supported FireVoxel software (wp.nyu.edu/firevoxel). Axial images were reformatting into sagittal (Fig. 2a) and coronal projections from which the slice corresponding to the superior endplate of the L3 vertebral body was identified. Quantitative adipose tissue analysis at L3 abdominal level was performed by a board certified abdominal radiologist with 12 years of the post-fellowship experience. The reader was blinded to the clinical outcome.
The semi-automated analysis includes the following steps:

1. Body masking was performed automatically to remove all outside objects unrelated to the abdomen such as the supporting table, patient arms, and tubing.
2. Automatic median filtering, followed by thresholding to the range of [-120, -90] Hounsfield Units (HU) and morphological noise removal yielded the total fat (TAT) mask (Fig. 2b).
3. The abdominal cavity contour was generated by placing between 10 and 20 anchor points (mouse clicks) along the margin of the peritoneal surface of the abdominal cavity (Fig. 2c).
4. TAT voxels were automatically partitioned in two classes: those inside the abdominal cavity contour were labeled as VAT, and those outside as SAT (Fig. 2d). The total areas of SAT (SATL3), VAT (VATL3), and TAT (TATL3) in cm² were computed.

To verify measurement repeatability, another observer (over 25 years of experience in image processing) independently calculated L3-level visceral and subcutaneous fat measures on a randomly selected subset of 25 cases as described above.

Fig. 1 Patient inclusion and exclusion criteria for study cohort of patients with PCR diagnosis of SARS-CoV-2 infection and abdominopelvic CT examination performed between March 19 and April 19, 2020.

Table 1 Demographics of patients with COVID-19

|                | Outpatient | Hospitalized but no MV | Hospitalized + MV |
|----------------|------------|------------------------|-------------------|
| Number of subjects | 10         | 30                     | 11                |
| Age            | 54.7 ± 11.6| 58.2 ± 17              | 67.9 ± 10.4       |
| Sex            | 5 M, 5 F   | 22 M, 8 F              | 11 M, 0 F         |
| BMI (kg/m²)    | 28.1 ± 7.6 | 30.4 ± 7.8             | 27.6 ± 2.3        |

After the step (3) in the analysis pipeline described above was performed on every fifth slice and automatically interpolated in z-direction, the remaining steps for computing VAT3D, and SAT3D were as for the single-slice analysis. All 3D measurements were done by a reader blinded to the clinical outcome as well as single-slice analysis.

**Statistical analysis**

Single-slice (at the L3 vertebral body level) VATL3, SATL3, TATL3, VAT/TATL3, and BMI were compared using Mann–Whitney test for independent samples between the outpatient and hospitalized patients, as well as in the hospitalized patients that did not require MV versus those that were admitted to the ICU and required MV (hospitalized + MV). These data were also stratified by gender. VATL3, SATL3, TATL3, and VAT/TATL3 measures were correlated with BMI. The intraclass correlation coefficient (absolute agreement version) was used to assess repeatability in VATL3, SATL3, and VAT/TATL3 in 25 subjects with measurement providers by two independent observers. Volumetric measure of VAT3D, SAT3D, and VAT/TAT3D were correlated to single-slice L3 measurements (in cm²) in 25 subjects.
In the next step, two logistic regression models were fit to discriminate patients requiring hospitalization from those that were outpatients. A clinical model that included age, sex, and BMI was compared to a clinical + CT model that also included VAT\textsubscript{L3}. The two models were compared with BIC and Chi-squared test. ROC analysis was performed and area under the curve (AUC) was computed for both of the models to discriminate patients requiring hospitalization from outpatients.

MedCalc (version 19.2.6) and R (version 3.6.3) were used to perform statistical analysis.

**Results**

Ten patients with a diagnosis of COVID-19 were not admitted to the hospital (outpatient), whereas the remaining 41 patients were hospitalized. Of these 41 hospitalized patients, 11 patients were admitted to the ICU and required MV, and the remaining 30 hospitalized patients were not admitted to the ICU.

**Age**

The mean age of all the patients was 59.8 ± 14.9 years. The age of the outpatients \((n = 10)\) was 54.7 ± 11.6 years and the age of all the hospitalized patients \((n = 41)\) was 60.8 ± 15.8 years. There was no significant difference between the outpatient and the hospitalized patients \((p > 0.05)\). Patients requiring MV and ICU admission were older with age of 67.9 ± 10.4 years. Although patients requiring MV and ICU admission were older compared to other hospitalized and outpatients this did not reach statistical significance.

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**Fig. 2**

a Selection of the axial slice at the superior end plate of L3 vertebral body. 

b Histogram of attenuation values within selected CT slice shows a well-defined peak with the maximum at −110 Hounsfield Unit (HU). 

c Axial slice with superimposed teal color overlay for voxels in [−120, −90] HU attenuation range. The expert observer partitioned the mask by placing a contour of approximately 20 anchor points (thin red line) to outline abdominal cavity. 

d Fat voxels inside the abdominal cavity contour were labeled as VAT (red) and fat voxels outside the abdominal cavity were identified as SAT (green).
Gender

There were a total of 13 women and 38 men. 50% of outpatients were men ($n = 5$), whereas 80% of all the hospitalized patients were men ($n = 33$). Furthermore, all the ICU patients were men ($n = 10$). This difference between the outpatient and hospitalized patients was statistically significant ($p = 0.047$).

BMI

The BMI in the outpatient and hospitalized patients were not significantly different (28.1 ± 7.6 kg/m² vs. 29.8 ± 6.7 kg/m²). Furthermore, there were no differences in BMI of the ICU patients (27.6 ± 2.3 kg/m²) versus hospitalized patients. All $p$ values > 0.05.

VATL3 and SATL3

The VATL3, SATL3, TATL3, and VAT/TATL3 in the three phenotypic cohorts based on disease severity are shown in Table 2. There was a significant difference in VATL3 and VAT/TATL3 in the hospitalized patients when compared to the outpatients (VATL3: 228.6 ± 111.1 cm² vs. 128.0 ± 92.1 cm², $p = 0.01$; and VAT/TATL3: 0.52 ± 0.14 vs. 0.35 ± 0.20, $p = 0.01$); Fig. 3. There were no significant differences in SATL3 and TATL3 between the hospitalized patients and the outpatients (all $p > 0.5$).

VATL3 in the patients on MV in the ICU was higher than in the hospitalized patients not requiring MV, but this was not statistically significant (Table 2). There were no significant differences in SATL3, TATL3, and VAT/TATL3 between patients requiring MV and hospitalized patients that did not require MV (all $p > 0.05$).

Comparison stratified by gender

Women: There were five female outpatients and eight hospitalized female patients. The VATL3 in hospitalized female patients was significantly higher than outpatients (175.8 ± 55.5 cm² vs. 92.4 ± 54.1 cm²; $p = 0.03$) (Table 3).

Men: There were five outpatient men and 33 hospitalized men. The mean VATL3, TATL3, and VAT/TATL3 were higher in hospitalized men compared to the outpatient men (Table 4), but these differences did not reach statistical significance (all $p > 0.05$).

Table 2 VATL3, SATL3, TATL3, and VAT/TATL3 in the three phenotypic cohorts based on disease severity

|           | Outpatient | Hospitalized but no MV | Hospitalized + MV |
|-----------|------------|------------------------|-------------------|
| VAT (cm²) | 128.0 ± 92.1 | 224.2 ± 115.9          | 240.6 ± 101.2     |
| SAT (cm²) | 232.3 ± 125.3 | 231.5 ± 142.2          | 179.6 ± 56        |
| TAT (cm²) | 360.4 ± 149.1 | 455.7 ± 201.0          | 420.2 ± 134.7     |
| VAT/TAT   | 0.35 ± 0.2   | 0.50 ± 0.16            | 0.56 ± 0.08       |

There was a significant difference in VATL3 and VAT/TATL3 in the hospitalized patients when compared to the outpatients ($p = 0.01$). There were no significant differences in SATL3 and TATL3 between the hospitalized patients and the outpatients (all $p > 0.5$).

Table 3 VATL3, SATL3, TATL3, and VAT/TATL3 in outpatients and hospitalized women

|           | Outpatient | Hospitalized | $p$ value |
|-----------|------------|--------------|-----------|
| VAT (cm²) | 92.4 ± 54.1 | 175.8 ± 55.5 | 0.03      |
| SAT (cm²) | 238.0 ± 83.6 | 308.8 ± 126.2 | > 0.05    |
| TAT (cm²) | 330.4 ± 81.7 | 484.6 ± 150.5 | > 0.05    |
| VAT/TAT   | 0.29 ± 0.15  | 0.39 ± 0.13   | > 0.05    |

Fig. 3a VATL3 was significantly higher in hospitalized patients compared to outpatients. b VAT/TATL3 was significantly higher in hospitalized patients compared to outpatients.
Correlation of BMI with VATL3 and SATL3

There was no significant correlation between VATL3 and BMI (0.21; \( p = 0.16 \)), whereas there was a statistically significant strong correlation between SATL3 and BMI (0.84; \( p < 0.001 \)) as well as TATL3 and BMI (0.72; \( p < 0.001 \)). There was inverse correlation between VAT/TATL3 and BMI (−0.39; \( p = 0.006 \)).

Repeatability of single-slice measurements

Intraclass coefficient correlation for VATL3, SATL3, and VAT/TATL3 was 0.99 for each of these measures. The average difference in VATL3 between the two readers was 1.8%, and it was 1.5% for SATL3.

Correlation between single-slice and volumetric measures

In 25 subjects, VATL3, SATL3, and VAT/TATL3 were correlated to volumetric measures. There were statistically strong correlation in VATL3-3D (0.87; \( p < 0.001 \)), SATL3-3D (0.96; \( p < 0.001 \)), and VAT/TATL3-3D (0.97; \( p < 0.001 \)).

Models to discriminate hospitalized patients from outpatients

Clinical (age, sex, BMI) and Clinical plus CT (includes VATL3) models were compared to identify patients that required hospitalization. The Clinical + CT model better fitted the data with BIC of 47.51 compared to the BIC of 50.39 for the clinical model alone. Chi-squared test demonstrated significant (\( p = 0.009 \)) improvement of clinical + CT model compared to the clinical model with value of 6.75. Area under the curve (AUC) of the clinical + CT was higher compared to the clinical model (0.847 versus 0.750) on ROC analysis for identifying patients requiring hospitalization (Fig. 4).

Discussion

A number of studies have highlighted an association of obesity with severity of COVID-19 symptoms, including hospitalization and need for invasive mechanical ventilation [7, 8]. It is well understood that there are different types of adipose tissue depots that contribute to obesity. These include VAT and SAT, which confer different degree of risks for metabolic disorders and cardiovascular risks [13, 15]. Furthermore, VAT is metabolically more active compared to SAT and is associated with release of pro-inflammatory markers [16]. Hence, in this study we assessed VATL3, SATL3, and TATL3 on abdominopelvic CT in SARS-CoV-2-positive patients with different severity of COVID-19 infection, and (2) assessed correlations between BMI and VATL3, SATL3, TATL3, and VAT/TATL3 estimates at CT in patients with COVID-19.

Our results demonstrated significant higher VATL3 and VAT/TATL3 in hospitalized patients compared to the outpatients. Patients admitted to ICU with need for MV had higher VATL3 compared to the hospitalized patients that were not admitted to ICU, but this was not statistically significant. It is important to note that many factors are important in assessing the need for MV including patient’s DNR (do not resuscitate) status or availability of the ICU beds. Hence, severity of disease is not the only factor contributing to the ICU admission and MV. This study did not explore these other variables. Men and women have different distribution

| Table 4 VATL3, SATL3, TATL3, and VAT/TATL3 in outpatients and hospitalized men |
|------------------|------------------|------------------|------------------|
|                  | Outpatient       | Hospitalized     | \( p \) value    |
| VAT (cm²)        | 163.6 ± 114.0    | 241.4 ± 138.5    | > 0.05           |
| SAT (cm²)        | 226.7 ± 168.2    | 195.5 ± 111.0    | > 0.05           |
| TAT (cm²)        | 390.3 ± 202.7    | 436.9 ± 211.0    | > 0.05           |
| VAT/TAT          | 0.42 ± 0.23      | 0.55 ± 0.18      | > 0.05           |

Fig. 4 ROC analysis demonstrates higher AUC of the Clinical + CT model (0.847) compared to the clinical model (0.750) in discriminating patients requiring hospitalization from the outpatients, and this difference was statistically significant.
of VAT and SAT, therefore, we stratified our results by gender. In female patients, VAT_L3 was significantly higher in hospitalized patients compared to outpatients. In men, VAT_L3 was also higher in hospitalized patients but this did not reach statistical significance.

BMI is a readily available clinical measure of obesity but it does not capture all dimensions of obesity, and adipose tissue distribution may provide additional information not captured in BMI [20–22]. It is interesting to note that BMI demonstrated significantly strong correlation with SAT_L3 and TAT_L3, but not with VAT_L3. Therefore, in these patients VAT_L3 seems to provide information which is not reflected in BMI. To test the hypothesis that CT assessment of VAT_L3 captured in BMI [20–22].

Including VAT_L3 to this model (Clinical + CT) significantly improved the performance of the model in discriminating outpatient from hospitalized patients with AUC of 0.847.

We performed single-slice evaluation of adipose tissue content at the L3 vertebral body level. A number of studies have shown that a single-slice evaluation provides good estimate of the total abdominal fat content [17]. To confirm this hypothesis in COVID-19 patients in our cohort we correlated VAT and SAT measures obtained on a single-slice to volumetric measures in 25 subjects. There was excellent statistically significant correlation between VAT_L3-3D, SAT_L3-3D, and VAT/TAT_L3-3D. While easier to integrate into clinical practice than volumetric measurements, single-slice workflow is semi-automated as it requires 1 to 2 minutes of user interaction. Our group is exploring an automated workflow solution for volumetric calculation of abdominal fat without any user interaction. Such a workflow could enable routine clinical use of volumetric measure of VAT, SAT, and VAT/TAT.

As in prior studies we also observed that the male gender was associated with hospitalization and ICU admission [7, 23] and when comparing outpatients to the hospitalized patients, this difference was statistically significant. Older age was also associated with hospitalization and ICU admission but in this small study these differences were not statistically significant. This was probably related to small sample size of our study.

Other risk factors such as diabetes mellitus, hypertension, history of cardiopulmonary disease, and immunocompromised state have also been identified as risk factors for hospitalization and severe disease [7, 23, 24]. In this small study we did not evaluate these other variables that are also associated with severe disease. This is one of the limitation of our study. Inclusion of these other variables in a larger study may further improve the ability to predict which patients at the time of presentation will require hospitalization. Furthermore, we did not separate intramuscular fat from SAT. The contribution of intramuscular fat can be assessed in the future as it could provide information about sarcopenia which could be an independent risk factor for hospitalization and severe outcomes. In this retrospective study, in some subset of outpatients the patient weight may have been self-reported rather than measured and this could impact BMI calculation.

In conclusion, this study demonstrates higher VAT was observed in COVID-19 patients that required hospitalization compared to the outpatients, and addition of VAT to the clinical model significantly improved its AUC in discriminating hospitalized from outpatients in this small study. More work needs to be done in larger datasets to assess if addition of VAT to clinical models can help stratify patient’s need for hospitalization and MV.

Author contributions All authors have fulfilled ICJME criteria for authorship.

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Code/software availability Firevoxel software is available upon request.

Compliance with ethical standards

Conflict of interest Research support not related to current project. Hersh Chandarana: Support in form of hardware and software from Siemens Healthineers.

Ethical approval IRB approved retrospective study.

Informed consent Waiver of consent.

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