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***Retrospective Cohort Study***

**Decreased cross-sectional muscle area in male patients with clear cell renal cell carcinoma and peritumoral collateral vessels**

Greco F *et al*. CcRCC collateral vessels and decreased TAMA

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**Abstract**

BACKGROUND

Sarcopenia is the loss of skeletal muscle mass (SMM) and is a sign of cancer cachexia. Patients with advanced renal cell carcinoma (RCC) may show cachexia.

AIM

To evaluate the amount of SMM in male clear cell RCC (ccRCC) patients with and without collateral vessels.

METHODS

In this study, we included a total of 124 male Caucasian patients divided into two groups: ccRCCa group without collateral vessels (*n* = 54) and ccRCCp group with collateral vessels (*n* = 70). Total abdominal muscle area (TAMA) was measured in both groups using a computed tomography imaging-based approach. TAMA measures were also corrected for age in order to rule out age-related effects.

RESULTS

There was a statistically significant difference between the two groups in terms of TAMA (*P* < 0.05) driven by a reduction in patients with peritumoral collateral vessels. The result was confirmed by repeating the analysis with values corrected for age (*P* < 0.05), indicating no age effect on our findings.

CONCLUSION

This study showed a decreased TAMA in ccRCC patients with peritumoral collateral vessels. The presence of peritumoral collateral vessels adjacent to ccRCC might be a fine diagnostic clue to sarcopenia.

**Key Words:** Cancer cachexia; Body composition; Clear cell renal cell carcinoma; Collateral vessels; Kidney cancer; Sarcopenia

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**Core Tip:** Clear cell renal cell carcinoma (ccRCC) can be detected with or without peritumoral collateral vessels. These vessels have been defined as enlarged capsular veins, stimulated by tumor-related effects. The presence of peritumoral collateral vessels around ccRCC is a poorly investigated phenomenon, with unclear clinical meaning. Here, we reported a novel association between peritumoral collateral vessels and loss of skeletal muscle in patients with ccRCC. The effect was not influenced by age, supporting the concept that peritumoral collateral vessels adjacent to ccRCC should drive clinicians’ attention towards cancer cachexia.

**INTRODUCTION**

Cancer cachexia is the reduction of adipose tissue and skeletal muscle (SM) which cannot be fully compensated with nutrition, resulting in progressive functional impairment[1]. This condition is due to energy disbalance during growth of the neoplasm[2]. Advanced neoplastic diseases can lead to loss of up to 85% of adipose and SM tissues[3]. Cancer cachexia and weight loss influence prognosis and response to therapy[4,5]. Renal cell carcinoma (RCC) patients with an advanced and metastatic disease are susceptible to cachexia. RCC patients have a relatively high prevalence of sarcopenia, the term for loss of SM mass (SMM)[4,6]. For example, sarcopenia was detected in up to 47% of patients with localised RCC and 29%-68% of patients with metastatic RCC[7-9]. Sarcopenic RCC patients have a worse overall survival than RCC patients without sarcopenia[10].

SM is not only part of the locomotor system but also produces and releases cytokines and myokines through the contraction of muscle fibres and thus has endocrine activity[11]. By releasing myokines into the circulation, SM can communicate with other organs such as adipose tissue, bone, the liver, and the brain, underlining the importance of this organ for regulating endocrine balance and decreasing risk of various diseases[12].

Body mass index (BMI) is an indicator used for obesity classification but does not convey information about body composition nor does it provide details about the quantity and distribution of different tissues such as SM and abdominal adipose tissue compartments. For this, computed tomography (CT) and magnetic resonance imaging (MRI) are gold standard methods for quantitative assessment and non-invasive tissue characterisation[13-19].

Peritumoural collateral vessels in RCC result from enlargement of capsular renal veins[19]. Gonadal vein recruitment can be present, especially in RCCs located at the lower renal pole[19]. Conversely, lesions located at the upper renal pole have different drainage routes including the adrenal and lower phrenic veins[19]. A study performed on 58 RCC patients reported 28 patients with peritumoural collateral vessels, of which 18 presented with gonadal vein recruitment[19]. Peritumoural collateral vessels with gonadal vein outflow were detected only in RCCs greater than 5 cm in diameter[19].

It is reasonable to speculate that increased blood demand due to tumour hypercellularity and neovascularisation, in possible association with main renal vein thrombosis, are factors contributing to the development of peritumoural collateral vessels in RCC patients. Hypercellularity could influence changes in cellular architecture leading to alternative routes of venous outflow that can become macroscopically evident as peritumoural collateral vessels with CT and MRI imaging (Figure 1). The presence of collateral vessels adjacent to RCC is considered a sign of locally advanced disease (*i.e.*, pT stage > T3a)[20]. However, these vessels can also be present in early stages of RCC.

The direct comparison of SMM in clear cell RCC (ccRCC) patients with and without peritumoural collateral vessels has not been performed to date. Evaluating the relationship between peritumoural collateral vessels in ccRCC patients and reductions of SMM would be of clinical interest for prognostic implications. We hypothesised that ccRCC patients would have a decreased cross-sectional total abdominal muscle area (TAMA) and peritumoural collateral vessels as a metabolic systemic consequence of locally advanced disease. To address this question, we evaluated SMM in male ccRCC patients with and without peritumoural collateral vessels using a CT imaging-based approach.

**MATERIALS AND METHODS**

This observational retrospective study was conducted in accordance with the Declaration of Helsinki. CT images and data from ccRCC patients with and without peritumoural collateral vessels were downloaded from the Cancer Imaging Archive (TCIA)[21-23]. This data collection received approval from our Institutional Review Board. The subsequent analysis contained publicly available and anonymised data which did not require further review due to previous protections implemented by TCIA. All enrolled subjects signed a written informed consent agreement.

A total of 267 patients with a histologically proven diagnosis of ccRCC were evaluated and selected by examining medical histories and CT images. The exclusion criteria for this study were: Female patients, patients with non-Caucasian ethnicity, patients who had undergone MRI examination only, patients who had undergone chest CT only, heminephrectomised and nephrectomised patients, patients with previous renal ablation, cirrhotic patients with collateral vessels, and patients with a congenital solitary kidney. The selected ccRCC patients were divided into two groups: Absence and presence of collateral vessels (ccRCCa and ccRCCp, respectively).

***CT analysis***

All ccRCC patients underwent CT examination. Horos v.4.0.0 RC2 software was used for acquisition of TAMA measurements with a semi-automatic function that allowed identification of SM tissue attenuation values (*i.e.,* range 10-40 Hounsfield units)[16]. TAMA (cm2) was defined as the sum of the areas of the abdominal muscles visible on an axial image located 3 cm above the lower margin of L3[16]. This area was measured by selecting a region of interest (ROI) on the following muscles: The rectus abdominis, transversus abdominis, external oblique, quadratus lumborum, iliocostalis lumborum, longissimus thoracis, spinalis thoracis, and psoas major[13]. All ROIs were independently drawn by two radiologists (F.G., 5 years of experience; C.A.M., 9 years of experience) who were blinded to the clinical data. The mean of the two measurements was utilised as the value for each subject.

***Statistical analysis***

Data distribution normality was assessed by the Shapiro-Wilk test. Comparison of TAMA between the ccRCCa and ccRCCp groups was performed using the Student’s *t*‐test. To rule out age-related effects, TAMA values were corrected by dividing individual values of TAMA by the age of each subject. Sub-analyses for TAMA assessment were performed by Student’s *t*‐tests between ccRCC patients with low (I/II) or high (III/IV) Fuhrman grade and between patients that were alive or deceased at the time of data collection. To evaluate the reliability of measurements by the two radiologists, the intraclass correlation coefficient for the TAMA measurements was calculated using Cronbach’s alpha (also known as coefficient alpha). Finally, Kaplan-Meier curves were included to assess survival of the ccRCCa and ccRCCp groups. The threshold of statistical significance was established at *P* < 0.05.

**RESULTS**

A total of 124 male Caucasian ccRCC patients were selected according to the exclusion criteria. The two groups were composed as follows: ccRCCa (*n* = 54; mean age: 57, range: 26-83) and ccRCCp (*n* = 70; mean age: 59.8, range: 34-84). The staging of ccRCCa group patients were as follows: 1 T1N0M0, 8 T1aN0M0, 21 T1aNxM0, 6 T1bN0M0, 7 T1bNxM0, 1 T1bNxM1, 3 T2N0M0, 1 T2NxM0, 2 T3aN0M0, 2 T3aN0M1, 1 T3bN0M0, and 1 T3bNxM0. The staging of ccRCCp group patients were as follows: 10 T1aNxM0, 1 T1aNxM0, 1 T1aN1M0, 2 T1bN0M0, 8 T1bNxM0, 5 T2N0M0, 1 T2N0M1, 4 T2NxM0, 2 T2aNxM0, 1 T2bN0M0, 9 T3aN0M0, 1 T3aN0M1, 5 T3aNxM0, 1 T3aN0M1, 8 T3aNxM1, 1 T3aN1M1, 3 T3bN0M0, 4 T3bNxM0, 1 T3bNxM1, 1 T4NxM0, and 1 T4N1M1.

Only three (2.41%) of 124 patients had renal vein thrombosis and these three were included in the ccRCCp group (4.28% of ccRCCp patients). No patients had segmental renal vein thrombosis. All patients of the ccRCCp group (*n* = 70; 100%) showed an exophytic growth pattern. In addition, 31.42% of ccRCCp patients had T1 stage (*n* = 22), 18.57% T2 (*n* = 13), 47.14% T3 (*n* = 33), and 2.85% T4 (*n* = 2). A total of 28 patients had a history of previous malignancy and 11 patients received a neoadjuvant treatment.

No significant difference was detected in the ages of the two groups (*P* = 0.21). A statistically significant difference between the ccRCCa and ccRCCp groups was obtained for TAMA (*P* < 0.05). These results are summarised in Table 1 and represented in Figure 2. Examples of CT cases showing the observed effect are shown in Figure 3. Statistically significant differences between the ccRCCa and ccRCCp groups were confirmed after TAMA values were corrected for age (*P* < 0.05) (Table 1).

No statistically significant differences (*P* = 0.66) were found between ccRCC patients with low (*n* = 44; 1 grade I and 43 grade II) and high (*n* = 80; 61 grade III and 19 grade IV) Fuhrman grades. These results are summarised in Table 2. Patients who were deceased (*n* = 33) at the time of data collection demonstrated a statistically significant reduction (*P* < 0.001) of TAMA in comparison to those that were still alive (*n* = 90) (Table 3). Cronbach’s alpha of the two tracers was 0.913, indicating excellent reliability. No significant differences in survival between the two groups (available data for 54 of 54 ccRCCa patients and 69 of 70 ccRCCp patients) were found based on the Kaplan-Meier method (log-rank test: *Z* = 1.88, *P* = 0.06) (Figure 4).

**DISCUSSION**

This study showed a significant decrease of SMM in the ccRCCp patient group compared to the ccRCCa group. Although SMM is expected to decrease with age, we did not find a significant difference between the ccRCCa and ccRCCp groups in terms of age. This finding was supported by analysis of age-corrected TAMA values. Since differences in SMM can segregate according to gender and ethnicity, only male Caucasian patients were included in the present study to eliminate these potentially confounding factors[24,25].

It has been hypothesised that contraction of myofibres can affect metabolism by triggering the release of humoral/exercise factors from SM which signal for an increase in glucose demand from distant organs[26]. The concept of humoral factors has been progressively developed since cytokine interleukin 6 (IL-6) was found to increase in response to physical exercise causing both autocrine and endocrine effects[27,28].

The cytokines and other peptides produced, expressed, and secreted by SM are called myokines. This term, suggested by Pederson *et al*[29], derives from the Greek words for "muscle" and "motion" and refers to such molecules that exert an endocrine effect on the human body. The physiological consequences of autocrine and paracrine action of myokines includes regulation of muscle growth and lipid metabolism. For example, the myokines produced during exercise, including IL-6, IL-7, IL-15, irisin, and leukaemia inhibitory factor, determine muscle growth by stimulating protein synthesis and hypertrophy. Conversely, myostatin, a member of the transforming growth factor β (TGF-β) superfamily, causes muscle atrophy[30,31]. Activin A, another member of TGF-β superfamily, reproduces the same action of myostatin on SM[30]. Increased blood levels of activin A is known to reduce muscle strength and has been positively correlated with cachexia in cancer patients[32].

Factors that can distort tumour extension such as peritumoural inflammation or the presence of a secondary pseudocapsule can reduce the effectiveness of CT in distinguishing T1 and T2 stages from T3a[19,33]. Incorrect staging, in fact, was detected in 27 of 94 tumours in a study of RCC patients using cross-sectional imaging[19]. Peritumoural collateral vessels in RCC patients showed a specificity of 94% and positive predictive value of 88% in staging of locally advance disease by cross-sectional CT imaging[19].

In our study, 100% of the patients from the ccRCCp group exhibited an exophytic growth pattern. This novel finding suggests a link between peritumour collateral vessels and the RCC growth pattern. Body composition imaging has gained an important role in the assessment of oncological risk, pathogenesis, and development of RCC[14-17]. CT imaging features of the tumour can also provide indications about the patient's body composition. In the present study, the peritumoural collateral vessels adjacent to the ccRCC was associated with a reduction of SMM, a possible sign of sarcopenia. Most likely, in ccRCCp patients, locally advanced disease determines muscle trophism loss as compared to ccRCCa patients. The progressive SMM reduction assessed by CT could be considered a sign of sarcopenia, and therefore of cancer cachexia, with potential prognostic implication for patients. Indeed, deceased ccRCC patients demonstrated a statistically significant reduction of TAMA relative to live patients, suggesting a link between sarcopenia and survival in our sample. However, Kaplan-Maier curves showed a difference just above the statistical threshold between the ccRCCa and ccRCCp patient groups.

The results of this study are supported by recent evidence showing a significant reduction of subcutaneous adipose tissue in ccRCC patients with peritumoural collateral vessels[17]. The limitations of this study include the retrospective study design which did not allow us to assess detailed clinical and anamnestic data including occupation, BMI, hormone blood levels, disease-free survival, timing of CT imaging, performing status, therapies, and CT follow-up after treatment. For instance, testosterone deficiency is known to be associated with an increase in proinflammatory cytokines. Inclusion of hormonal data, such as testosterone levels, could help better understand the cytokine cascade that is associated with pathogenesis and changes in body composition[34,35]. Similarly, CT follow-up after treatment (*e.g.,* surgery or chemotherapy/targeted immunotherapy) would have been helpful to understand changes in the sarcopenia index and the relationship with peritumoural collateral vessels after treatment. The vendor, model, and acquisition parameters (such as slice thickness) of the CT imaging used in this study were also unavailable. Images from the open-source TCIA were often acquired heterogeneously at multiple centres as part of clinical routine. A larger sample size would have strengthened our multivariate assessment of whether collateral vessels are an independent predictor of sarcopenia as well as the potential impact of other variables such as staging[36-38].

Further studies are needed to evaluate sarcopenia index changes after treatment to add robustness to the role of peritumoural collateral vessels as a prognostic biomarker for ccRCC patients. Such studies should consider abdominal circumference and patients’ occupation, which is a factor that can influence SMM (for example, people who are engaged in heavy physical labour would be expected to have significantly more muscle mass compared to office workers)[39]. Finally, SMM content of other subtypes of kidney cancer (*e.g.,* chromophobe and papillary) or other categories of cancer patients should be evaluated to assess the impact of SMM trophism on a patient's health status and prognosis.

**CONCLUSION**

This study showed a reduction of SMM in ccRCC patients with peritumoural collateral vessels. The presence of peritumoural collateral vessels adjacent to ccRCC is a good candidate biomarker for sarcopenia and therefore of cancer cachexia.

**ARTICLE HIGHLIGHTS**

***Research background***

Sarcopenia is the loss of skeletal muscle mass (SMM) and is part of cancer cachexia in which there is a decrease of adipose tissue and SM. Peritumoral collateral vessels adjacent to renal cell carcinoma (RCC) are indicative of locally advanced disease.

***Research motivation***

Metabolic systemic consequence related to a locally advanced disease might be linked to a decrease of SSM in clear cell RCC (ccRCC) patients with peritumoral collateral vessels, possibly providing clinically relevant information.

***Research objectives***

The aim of this study was to evaluate the amount of SMM in male ccRCC patients with and without peritumoral collateral vessels, in order to understand a possible relationship between sarcopenia and collateral vessels.

***Research methods***

In this study, we included a total of 124 male Caucasian patients divided into two groups: ccRCCa (*n* = 54) and ccRCCp (*n* = 70) groups, respectively, without and with collateral vessels. Computed tomography imaging-based approach was used for total abdominal muscle area (TAMA) measurements.

***Research results***

There was a statistically significant difference between the two groups for TAMA (*P* < 0.05).

***Research conclusions***

This study showed a reduction of TAMA in male ccRCC patients with peritumoral collateral vessels.

***Research perspectives***

Further studies, on larger sample size and with longitudinal data, will shed light on collateral vessels adjacent to RCC as a possible biomarker of cachexia and sarcopenia.

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**Footnotes**

**Institutional review board statement:** All the procedures were retrospective and agreed with the Declaration of Helsinki. CT images and data of ccRCC patients were retrieved from The Cancer Imaging Archive (TCIA). The TCIA project received approval of the Institutional Review Board. This subsequent retrospective analysis was on the publicly available, anonymized data and did not require further review due to previous protections implemented by TCIA.

**Conflict-of-interest statement:** The authors declare that they have no conflict of interest to disclose.

**Data sharing statement:** The data presented in this study are openly available in The Cancer Imaging Archive (<https://wiki.cancerimagingarchive.net/display/Public/TCGA-KIRC>, accessed on 1 November 2019).

**STROBE statement:** The authors have read the STROBE Statement—checklist of items, and the manuscript was prepared and revised according to the STROBE Statement—checklist of items.

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**Figure Legends**



**Figure 1 Axial computed tomography image shows the presence of clear cell renal cell carcinoma collateral vessels with the typical tortuous course located in the retroperitoneal space (arrow).**



**Figure 2 Bar chart with error bars showing a significant difference in mean values of total abdominal muscle area between the two groups.** ccRCC: Clear cell renal cell carcinoma.



**Figure 3 Axial computed tomography images with maximum intensity projection reconstruction of an 84-year-old male clear cell renal cell carcinoma patient without collateral vessels and an 82-year-old male clear cell renal cell carcinoma patient with collateral vessels.** These images show skeletal muscle masses (SMMs) and tumors in a clear cell renal cell carcinoma patient without collateral vessels (ccRCCa) (A) and a clear cell renal cell carcinoma patient with collateral vessels (ccRCCp) (B) (orange and dark orange arrows in A and B, respectively), as well as collateral vessels adjacent to the tumor in the ccRCCp patient (light blue arrows in B) and nodal metastasis infiltrating the ureter (yellow arrows in B). Please note the decrease of SMM clearly evident in the ccRCCp patient (B) compared to the ccRCCa patient (A).



**Figure 4 Kaplan-Meier curves showing no statistically significant difference of survival between the two groups (ccRCCa group is depicted as blue curve and ccRCCp group is depicted as red curve).**

**Table 1** **Total abdominal muscle area and total abdominal muscle area corrected for age in the two groups**

|  |  |  |
| --- | --- | --- |
|  | **TAMA (cm2)** | **TAMA\_C (cm2)** |
| ccRCCa group (mean, range, and SD) | 164.02 (91, 233.5 ± 31.86) | 3.08 (1.29, 5.83 ± 1.06) |
| ccRCCp group (mean, range, and standard deviation) | 150.91 (76.3, 218.3 ± 30.34) | 2.67 (1, 4.67 ± 0.91) |
| *P* | 0.02 | 0.02 |

TAMA: Total abdominal muscle area; TAMA\_C: Total abdominal muscle area corrected for age; ccRCC: Clear cell renal cell carcinoma.

**Table 2 Total abdominal muscle area of clear cell renal cell carcinoma patients with low Fuhrman grade (I/II) and high Fuhrman grade (III/IV)**

|  |  |
| --- | --- |
|  |  **TAMA (cm2)** |
| ccRCC patients with low Fuhrman grade (I/II) (mean, range, and standard deviation) | 158.27 (83.2-233.5), 35.41 |
| ccRCC patients with high Fuhrman grade (III/IV) (mean, range, and standard deviation) | 155.71 (76.3-219.2), 29.44 |
| *P* | 0.66 |

TAMA: Total abdominal muscle area; ccRCC: Clear cell renal cell carcinoma.

**Table 3 Total abdominal muscle area of alive and dead clear cell renal cell carcinoma patients**

|  |  |
| --- | --- |
|  | **TAMA (cm2)** |
| Alive ccRCC patients | 162.02 |
| (mean, range, and standard deviation) |  91, 233.5 ± 28.42 |
| Dead ccRCC patients | 150.91 |
| (mean, range, and standard deviation) | 76.3, 219.2 ± 34.84 |
| *P* | 0.0008 |

TAMA: Total abdominal muscle area; ccRCC: Clear cell renal cell carcinoma.



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