Computer vision applied to dual-energy computed tomography images for precise calcinosis cutis quantification in patients with systemic sclerosis

Anita C. Chandrasekaran 1, Zhicheng Fu 3, Reid Kraniski 4, F. Perry Wilson 5, Shannon Teaw 1, Michelle Cheng 1, Annie Wang 4, Shangping Ren 2,6, Imran M. Omar 7 and Monique E. Hinchcliff 1,5,8*

Abstract

Background: Although treatments have been proposed for calcinosis cutis (CC) in patients with systemic sclerosis (SSc), a standardized and validated method for CC burden quantification is necessary to enable valid clinical trials. We tested the hypothesis that computer vision applied to dual-energy computed tomography (DECT) finger images is a useful approach for precise and accurate CC quantification in SSc patients.

Methods: De-identified 2-dimensional (2D) DECT images from SSc patients with clinically evident lesser finger CC lesions were obtained. An expert musculoskeletal radiologist confirmed accurate manual segmentation (subtraction) of the phalanges for each image as a gold standard, and a U-Net Convolutional Neural Network (CNN) computer vision model for segmentation of healthy phalanges was developed and tested. A validation study was performed in an independent dataset whereby two independent radiologists manually measured the longest length and perpendicular short axis of each lesion and then calculated an estimated area by assuming the lesion was elliptical using the formula long axis/2 × short axis/2 × π, and a computer scientist used a region growing technique to calculate the area of CC lesions. Spearman’s correlation coefficient, Lin’s concordance correlation coefficient with 95% confidence intervals (CI), and a Bland-Altman plot (Stata V 15.1, College Station, TX) were used to test for equivalence between the radiologists’ and the CNN algorithm-generated area estimates.

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Background
Calcinosis cutis (CC) is defined as the deposition of insoluble calcium salts within the skin and subcutaneous tissues [1]. There are five subtypes of calcinosis cutis: dystrophic, metastatic, iatrogenic, calciphylaxis, and idiopathic [1]. Dystrophic calcification from local tissue injury is the most common subtype and is associated with autoimmune disorders including systemic sclerosis (SSc) and juvenile dermatomyositis (JDM). The clinical presentation is highly variable including an incidental finding on imaging, lesions in the hands or pressure point areas resulting in soft tissue swelling and ulceration, to disabling and life-threatening lesions [1]. Upwards of 18–49% of SSc patients [1], 40% of JDM patients [2], and 1% of chronic renal dialysis patients [3] experience CC.

Calcinosis cutis remains a therapeutic challenge. Medications for CC include calcium channel blockers (e.g., diltiazem), warfarin, topical and intravenous sodium thiosulfate and aluminum hydroxide (chelation therapy), ceftriaxone, minocycline, colchicine, intravenous immunoglobulin (IVIg), and probenecid among others [1]. The data supporting the efficacy of these treatments are primarily from case reports, case series, and retrospective studies rather than randomized clinical trials [1]. Valid clinical trial design to test the safety and efficacy of proposed systemic treatments requires a feasible, precise, and reproducible method for CC burden quantification.

Dual-energy computed tomography (DECT), also known as “spectral imaging” [4], is an imaging modality established for the detection of coronary calcifications [5] and uric acid nephrolithiasis [6], and has more recently been studied for the identification and quantification of soft tissue monosodium urate deposits in patients with gout [7]. Single-energy computed tomography (CT) uses a single polychromatic X-ray beam, which is emitted and received from a single source and detector, respectively [8]. This limits its use in the differentiation of materials (e.g., fat, soft tissues) that have similar linear attenuation coefficients [8]. In contrast, DECT utilizes two energy levels [(usually 80 and 140 kilovoltage peaks (kVp)] [4], that results in greater differentiation between attenuation coefficients [9] and is particularly efficient for the evaluation of small body parts including the extremities [8]. Dual-energy computed tomography permits differentiation of CC lesions from adjacent bone, but radiologists must manually quantify lesional CC areas, which is time-consuming and therefore costly, and prone to inter-rater variability. Moreover, CC lesions are often irregularly shaped, which can lead to imprecise and inaccurate measurements. By convention, CC lesions are measured in three planes, using the maximal dimension as the reference axis, with two additional orthogonal axes. Using the ellipsoid formula, utilized in other clinical applications such as prostate measurement, a multiplication factor of 0.52 can then be incorporated for volume estimation [10, 11]. On subsequent exams, lesions need to be measured using the same axes, to maintain consistency and accuracy.

Computer vision, a field in which computers are trained to independently identify and process images [12], has been increasingly successfully applied to solve problems in clinical medicine. Since the late 1990s, computer scientists have been using large quantities of curated images to develop software systems that enable computers to perform tasks that previously required an
expert such as skin disease diagnoses [13–15]. Deep learning methods (e.g., convolutional neural networks [CNN], deep neural networks [DNN], and support vector machines [SVM]) are a more complex form of machine learning and have been developed to enable computers to filter inputs, such as images, sequentially through layers in order to learn how to predict and classify information without manual feature extraction (Fig. 1a) [12]. The process of filtering images using layers whereby the output of one layer becomes the input of a subsequent layer is the same process that neurons in the human brain use for image classification (e.g., car or clock), hence deep learning models are called neural networks [16].

Because large curated training data sets are not available for most diseases, a process called transfer learning was developed to enable algorithms that were developed for a disease with high prevalence to be applied to more rare diseases like SSc-CC [17]. Herein, we optimized a deep learning model using a CNN approach and applied this technology to DECT images of fingers from patients with SSc and clinically evident CC. The goal was to train a computer to quantify CC area lesions that currently requires a highly trained musculoskeletal radiologist. The use of artificial intelligence for accurate, precise, and rapid quantification of CC burden in patients with SSc will facilitate musculoskeletal radiologists’ workflow and improve clinical trial design.

**Methods**

**Identification of DECT images containing calcinosis cutis in patients with systemic sclerosis**

We conducted a retrospective study in patients with SSc and clinically evident finger CC lesions who were referred for 2-dimensional (2D) DECT imaging. Because no personal health information was collected and all images were de-identified, the study was not considered human subjects research and was exempt from institutional board approval. The Northwestern Memorial Hospital electronic health record was queried for patients with a diagnosis of SSc (ICD9 = 710.1 or ICD10 = M34.9) and DECT hand images. De-identified sagittal reformatted images from patients were downloaded from the Picture Archiving and Communication System (PACS) in jpeg form to a secure server maintained by the Northwestern University Information Technology, then

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**Fig. 1** a Distinction between machine learning and deep learning. b Optimization of the CNN algorithm for forefinger bone segmentation
transferred to Visage Imaging 7 (San Diego, CA) for examination purposes. All images were optimized to assess bone (window of 1500 and level of 450). Because bone and CC lesions both consist of basic calcium hydroxyapatite, a computer scientist manually segmented (identified) finger phalanges on training set images and an expert musculoskeletal radiologist (IMO) confirmed measurements that were then used as a gold standard reference (ground truth) for the deep learning models (Fig. 1).

CNN software training
Various segmentation algorithms (e.g., expectation maximization [18], Fuzzy c-means [19], K-means [20], mean-shift segmentation [21], V-Net [22], P-Net [23], and U-Net [24]) were tested in order to identify the optimal approach for finger bone segmentation. U-Net CNN performed best and was utilized for the experiments herein.

Training set and test set one
Forty sagittal reformatted DECT finger images from three unique patients were divided into a training set (N = 30) and a test set (N = 10). To enlarge the training dataset to permit development of a U-Net CNN model, 30 images were spatially rotated three times for a total of 90 additional images (total 120 images). A NVIDIA Tesla k40c GPU computing processor with varying numbers of epochs was used (Fig. 1). Visual inspection was conducted between runs to determine the concordance between manual and computer bone segmentation in order to identify the optimal number of epochs. The segmentation approach was tested in ten independent images and manual inspection confirmed segmentation accuracy.

Calcinosis cutis area estimates
Thirty-one sagittal reformatted DECT finger images containing CC from 13 unique patients were identified as an independent test. The CNN finger bone segmentation algorithm was applied, arithmetic operations were applied to segmented images to subtract out the normal phalanges, and region growing, a pixel-based image segmentation method, was deployed to quantify the area of residual CC lesions [25]. Concurrently, a musculoskeletal radiologist (IMO) measured each lesion’s maximal long axis and maximal perpendicular short axis (mm) and then calculated an estimated area of each lesion by assuming the lesion was elliptical and using the formula long axis/2 × short axis/2 × π. A second expert musculoskeletal radiologist (AW) independently measured the area of each CC lesion, and the concordance between radiologists’ measurements was assessed.

Statistics
First, CC lesional area estimates were compared between radiologists (IMO and AW) to test the precision of area estimates. Next, the radiologist (IMO)- and CNN-generated CC area estimates were compared to test the accuracy of the CNN approach. Spearman’s correlation coefficient was used to assess the correlation, and Lin’s concordance correlation coefficient (CCC) was used to test for equivalence, while Bland-Altman plots were used to evaluate agreement. A p value ≤ 0.05 was considered significant. Stata V 15.1, College Station, TX, was used for analyses.

Results
Training the CNN model
The training set (30 images each rotated three times to generate 120 images) was used to train the segmentation algorithm. One hundred, 200, 300, 400, and 500 epochs (iterations) were evaluated and by manual inspection, and 500 epochs was deemed the optimal number (Figs. 1b and 2). A 4-h CNN run time was required for the CNN to learn to discriminate healthy finger bones from adjacent CC lesions (Fig. 3). The algorithm was tested in ten additional DECT finger images, and manual inspection was used to confirm segmentation accuracy.

Measurement of calcinosis cutis lesions in an independent test set
Next, the accuracy of the computer vision approach for estimating the CC area was determined in an independent set of 31 images (test set two), by comparing the area estimates generated by computer vision (region growing technique) to those generated by an expert musculoskeletal radiologist. The U-Net CNN algorithm was unable to be applied to seven out of 31 (23%) test set images due to incomplete visualization of the phalanges (Fig. 4). For the 31 images scored by both radiologists and computer vision, the area range for radiologist (IMO) was 0.5–41.5 mm² and for radiologist (AW) was 0.4–40.7 mm². Per the CNN algorithm, 2D area ranged from 0.69–48.18 mm².

The correlation between the two radiologists’ measurements was excellent (Spearman’s rho 0.91, p < 0.0001, Lin’s concordance correlation coefficient 0.91 [95% CI 0.85–0.98, p < 0.001] (Fig. 5a) and a Bland-Altman plot demonstrating a mean difference −0.5 mm² (95% limits of agreement −10.0–9.0 mm²) (Fig. 5b). Similarly, the correlation between the radiologist vs. computer vision area estimates was also excellent (Spearman’s rho 0.94, p < 0.001), Lin’s concordance correlation coefficient 0.95 [95% CI 0.91–0.99, p < 0.001] (Fig. 5c), and a Bland-Altman plot demonstrating a mean difference −1.7 mm² (95% limits of agreement −6.0–9.5 mm²) (Fig. 5d).
Discussion
Calcinosis cutis can significantly worsen health-related quality of life, depending on lesion location and size, increasing the risk of infection, and leading to deformities, particularly in patients with SSc [1]. While deep learning technological advancements have improved the performance of image recognition in the field of medical image analysis including the diagnosis of skin diseases [26], computer vision has not been previously applied to the problem of CC quantification. Given a lack of standardized and validated methods to assess CC disease burden, accurate and precise CC quantification represents an unmet medical need. This study evaluated the utility of applying a CNN algorithm to DECT hand images of SSc patients to differentiate CC lesions from adjacent healthy bone and to permit rapid and reproducible quantification of CC lesional area. We found that the computer vision method applied to finger images had a substantial degree of concordance and was highly accurate compared to gold standard radiologist area estimates. These data suggest that computer vision may be a useful approach for CC disease burden quantification.
The difficulty of assessing CC severity in SSc patients has been addressed in prior studies. Utilizing plain X-rays, Chung et al. in 2014 aimed to develop and validate an original scoring system for CC affecting the hands of SSc patients [27]. They developed a feasible scoring system that accounted for lesional area, density, and anatomic location, but significant interrater and intra-rater reliability were noted [27]. In the present study, we used DECT imaging of the hands because this technique has been successfully used to identify and quantify soft tissue monosodium urate deposits in patients with gout [9]. Thus, we hypothesized that DECT would be a useful approach for imaging and quantifying CC in the hands of patients with SSc.

Conventional imaging studies, such as computed tomography (CT) scans and magnetic resonance imaging (MRI), have not been proven to be reliable in the identification of small, irregular calcifications. Newer diagnostic techniques, such as ultrasonography (US), multidetector computed tomography (MDCT), and DECT, have been evaluated in small trials for CC detection [28–31]. A case-control study using US in 44 patients with SSc demonstrated the presence of CC in 17 patients (39%), with a sensitivity of 89% compared to radiography [30]. Multidetector computed tomography has higher tissue resolution and generates 3D images but requires higher radiation exposure and has not been studied in CC [28]. There are minimal current data on the use of DECT in CC. A small study in 2015 of 16 patients with SSc-CC in the hands found that DECT imaging was superior to plain radiographs in localizing soft tissue CC though CC lesions and bone were noted to have the same color and density [7].

We examined the utility of applying CNN, specifically U-Net, to the problem of CC measurement in SSc because U-Net was initially developed for biomedical image segmentation [32]. The two main benefits of U-Net are (1) the ability to develop a useful model in spite of a small training dataset and (2) the U-architecture and skip-connection performance performs better for segmenting different levels of semantic information. The first task to train the computer to “see” the finger bones was readily accomplished in healthy finger bones from SSc patients with normal anatomy. The algorithm failed to recognize the finger bones in cases where the CC lesion was in an area where the entire phalange was not visible (Fig. 4).

Our study has several strengths. Convolutional neural networks have recently been applied for cartilage segmentation on knee MRI scans [33], brain lesion segmentation [34], automatic polyp detection on colonoscopy videos [35], and pulmonary embolism detection on CT [36], but to our knowledge, this is the first study to apply deep learning to analyze CC lesions in SSc patients. Our
study involved two expert musculoskeletal radiologists who independently measured CC lesional areas to permit interrater reliability. The computer algorithm was trained to segment the phalanges assessment using real 2D DECT images from SSc patients with clinically evident lesser finger CC lesions and demonstrated a high degree of concordance with a radiologist’s gold standard assessment, despite variable lesional shape and size. This suggests that this deep learning model may be an efficient and reproducible method for CC measurement.

Several limitations are noted as well. This study focused on the hands, specifically lesser finger lesions, because the hands/fingers have been shown to be the area most frequently affected by CC in SSc patients [37]. To be useful in clinical trials, the CNN algorithm will need to be optimized and tested in other anatomic locations. The algorithm can only quantify the area of CC lesions when all three phalanges are visible, limiting its usefulness (Fig. 4). Finally, the principles of 2D segmentation can be transferred to 3-dimensional (3D) segmentation but does require increased computational complexity that was not performed in this study. V-Net is an example of a CNN utilized for 3D segmentation, previously applied to prostate gland segmentation on MRI scans [22], that may be used in future work for CC density quantification, crucial for assessing response to treatment.

Conclusions
In this study, we demonstrate that computer vision approaches can be applied to the problem of CC quantification in patients with SSc. Expert musculoskeletal radiologist review may not be locally available at many centers, and thus, the ability to add a CNN tool into standard radiology software programs such as the Visage Imaging 7 platform might be advantageous. Convolutional neural network-derived CC quantification had a high degree of correlation with expert radiologist measurement. Moreover, our approach may be generalizable to other diseases that cause dystrophic calcification, such as chronic kidney disease, JDM, and malignancy. To facilitate optimization of computer vision approaches to aid in the quantification of CC, a de-identified dataset of our DECT images has been made publicly available online (https://drive.google.com/drive/folders/1_tXg3dnSQQrZ4_Ro-0JZ3h_hMv0H1e9h?usp=sharing). Future
work will involve transferring the principles of 2D segmentation to 3D segmentation for volumetric and density CC quantification as well as additional validation in larger, multi-center SSc cohorts.

Abbreviations
CC: Calcinsis cutis; SSc: Systemic sclerosis; JDM: Juvenile dermatomyositis; IVIG: Intravenous immunoglobulin; DECT: Dual energy computed tomography; CT: Computed tomography; CNN: Convolutional neural network; SVM: Support vector machine; 2D: Two-dimensional; PACS: Picture Archiving and Communication System; CCC: Concordance correlation coefficient; SD: Standard deviation; US: Ultrasound; MDCT: Multidetector-computed tomography; 3D: Three-dimensional

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Authors’ contributions
MH, IMO, and SR designed the study. MH, IMO, AW, RK, AC, ST, MC, FPW, and ZC analyzed the data. FPW performed statistical analyses. AC and MH wrote the manuscript. The authors read and approved the final manuscript.

Authors’ information
FPW and MH are founding members of the Clinical and Translational Research Accelerator (CTRA) at the Yale School of Medicine. Shangping Ren, PhD, is the Chair of the Department of Computer Science at San Diego State University and Zhicheng Fu, PhD, currently works at Motorola, Inc.

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Availability of data and materials
The datasets generated and/or analyzed during the current study will be made publicly available.

Ethics approval and consent to participate
The Northwestern University Institutional Review Board declared this study exempt because all images are de-identified and no personal health information was studied.

Consent for publication
Not applicable

Competing interests
MH has received consulting fees from Boehringer Ingelheim and AbbVie.

Author details
1. Yale School of Medicine, Section of Rheumatology, Allergy & Immunology, The Alynn Center, 300 Cedar Street, PO BOX 208031, New Haven, CT 06520, USA. 2. Department of Computer Science, Illinois Institute of Technology, 10 W 31st St, Chicago, IL 60616, USA. 3. Motorola Mobility LLC, 222 W Merchandise Mart Plaza #1800, Chicago, IL 60654, USA. 4. Department of Radiology, Yale School of Medicine, 330 Cedar St, New Haven, CT 06520, USA. 5. Clinical and Translational Research Accelerator, Department of Medicine, Yale School of Medicine, Temple Medical Center, 60 Temple Street Suite 6C, New Haven, CT 06510, USA. 6. Department of Computer Science, San Diego State University, 5500 Campanile Drive, San Diego, CA 92182, USA. 7. Department of Radiology, Northwestern University Feinberg School of Medicine, 676 N St Clair St, Chicago, IL 60611, USA. 8. Department of Medicine, Division of Rheumatology, Northwestern University Feinberg School of Medicine, 240 E Huron Street, Suite M-300, Chicago, IL 60611, USA.

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References
1. Valenzuela A, Song P, Chung L. Calcinsis in scleroderma. Curr Opin Rheumatol. 2018;30(6):554–61.
2. Hoeltzel MF, Oberle EJ, Robinson AB, Agarwal A, Rider LG. The presentation, assessment, pathogenesis, and treatment of calcinosis in juvenile dermatomyositis. Curr Rheumatol Rep. 2014;16(12):467.
3. Rivet J, Lebbe C, Urena P, Cordoliani F, Martinez F, Baglin AC, et al. Cutaneous calcification in patients with end-stage renal disease: a regulated process associated with in situ osteopontin expression. Arch Dermatol. 2006;142(7):900–6.
4. Grajo JR, Patino M, Prochowski A, Sahani DV. Dual energy CT in practice: basic principles and applications. Appl Radiol. 2016;45(7):12–6.
5. Reimann AJ, Rinck D, Brinici-Aygadan A, Scheuering M, Burgstahler C, Schroeder S, et al. Dual-source computed tomography: advances of improved temporal resolution in coronary plaque imaging. Invest Radiol. 2007;42(3):196–203.
6. Graser A, Johnson TR, Bader M, Staehler M, Haske N, Nikolau K, et al. Dual energy CT characterization of urinary calciﬁ: initial in vitro and clinical experience. Invest Radiol. 2008;43(2):112–9.
7. Hsu V, Bramwit M, Schlesinger N. Use of dual-energy computed tomography for the evaluation of calcinosis in patients with systemic sclerosis. Clin Rheumatol. 2015;34(9):1557–61.
8. Goo HW, Goo JM. Dual-energy CT: New horizon in medical imaging. Korean J Radiol. 2017;18(4):555–69.
9. Chow H, Chin TY, Peh WC. Dual-energy CT in gout - a review of current concepts and applications. J Med Radiat Sci. 2017;64(1):41–51.
10. Garvey B, Turkbey B, Troong H, Bernardo M, Peitarsawmy S, Choyke PL. Clinical value of prostate segmentation and volume determination on MRI in benign prostatic hyperplasia. Diagn Interv Radiol. 2014;20(3):229–33.
11. Breau RH, Clark E, Bruner B, Cervi P, Aswell T, Knoll G, et al. A simple method to estimate renal volume from computed tomography. Can Urol Assoc J. 2013;7(5–6):189–92.
12. Litjens G, Kooi T, Bejnordi BE, Setio AAT, Ciompi F, Ghafoorian M, et al. A survey on deep learning in medical image analysis. Med Image Anal. 2017;42:60–88.
13. Esteva A, Kuprel B, Novoa RA, Ko J, Swetter SM, Blau HM, et al. Dermatologist-level classification of skin cancer with deep neural networks. Nature. 2017;542(7639):115–8.
14. Olsen TG, Jackson BH, Feesper TA, Kent MN, Moad JC, Krishnamurthy S, et al. Diagnostic performance of deep learning algorithms applied to three common diagnoses in dermatopathology. J Pathol Inform. 2019;10:32.
15. Pal A, Gara R, Chandla A, Chatterjee R, Senapati S. Prioriasis skin biopsy image segmentation using deep convolutional neural network. Comput Methods Prog Biomed. 2018;159:59–69.
16. Marblestone AH, Wayne G, Kording KP. Toward an integration of deep learning and neuroscience. Front Comput Neurosci. 2016;10:94.
17. Correia C, Mawe S, Lofgren S, Marangoni RG, Lee J, Saber R, et al. High-throughput quantitative histology in systemic sclerosis skin disease using computer vision. Arthritis Res Ther. 2020;22(1):48.
18. Canin C, Belongie S, Greenspan H, Mall J. Bia image: segmentation using expectation-maximization and its application to image querying. IEEE Trans Pattern Anal Mach Intell. 2002;24(8):1026–38.
19. Zhong-dong W, Wei-xin X, Jian-ying Y, editors. Fuzzy C-means clustering algorithm based on kernel method. Proceedings Fifth International Conference on Computational Intelligence and Multimedia Applications (ICICMA 2003, 2003 27–30 Sept. 2003.
20. Dhanachandra N, Manglem K, Chau YJ. Image segmentation using K-means clustering algorithm and subtractive clustering algorithm. Procedia Computer Science. 2015;54:764–71.
21. Comaniciu D, Meer P. Mean shift; a robust approach toward feature space analysis. IEEE Trans Pattern Anal Mach Intell. 2002;24(5):603–19.
22. Milletarì F, Navab N, Ahmadî SA, editors. V-net: fully convolutional neural networks for volumetric medical image segmentation. 2016 Fourth International Conference on 3D Vision (3DV); 2016: IEEE.
23. Wang G, Zuluaga MA, Li W, Pratt R, Patel PA, Aertsens M, et al. DeepGeoS: a deep interactive geodesic framework for medical image segmentation. IEEE Trans Pattern Anal Mach Intell. 2019;41(7):1559–72.
24. Ronneberger O, Fischer P, Brox T, editors. U-net: convolutional networks for biomedical image segmentation. International Conference on Medical image computing and computer-assisted intervention; 2015: Springer.

25. Fan J, You DY, Elmagarmid AK, Aref WG. Automatic image segmentation by integrating color-edge extraction and seeded region growing. IEEE Trans Image Process. 2001;10(10):1454–66.

26. Liu Y, Jain A, Eng C, Way DH, Lee K, Bui P, et al. A deep learning system for differential diagnosis of skin diseases. Nat Med. 2020;26(6):900–8.

27. Chung L, Valenzuela A, Fiorentino D, Stevens K, Li S, Harris J, et al. Validation of a novel radiographic scoring system for calcinosis affecting the hands of patients with systemic sclerosis. Arthritis Care Research. 2015;67(3):425–30.

28. Freire V, Becce F, Feydy A, Guerini H, Campagna R, Allanore Y, et al. MDCT imaging of calcinosis in systemic sclerosis. Clin Radiol. 2013;68(3):302–9.

29. Elhai M, Guerini H, Bazel R, Avouac J, Freire V, Drape JL, et al. Ultrasonographic hand features in systemic sclerosis and correlates with clinical, biologic, and radiographic findings. Arthritis Care Research. 2012; 64(8):1244–5.

30. Freire V, Bazel R, Elhai M, Campagna R, Pessis E, Avouac J, et al. Hand and wrist involvement in systemic sclerosis: US features. Radiology. 2013;269(3):824–30.

31. Hsu V, Bramwit M, Schlesinger N. Dual-energy computed tomography for the evaluation of calcinosis in systemic sclerosis. J Rheumatol. 2015;42(2):345–6.

32. Ronneberger O, Fischer P, Brox T. U-Net: convolutional networks for biomedical image segmentation. CoRR. 2015;abs/1505.04597.

33. Prasoon A, Petersen K, Igel C, Lauze F, Dam E, Nielsen M. Deep feature learning for knee cartilage segmentation using a triplanar convolutional neural network. Med Image Comput Comput Assist Interv. 2013;16(Pt 2):246–53.

34. Kamnitas K, LEDIG C, NEWCOMBE VFJ, SIMPSON JP, KANE AD, MONO DK, et al. Efficient multi-scale 3D CNN with fully connected CRF for accurate brain lesion segmentation. Med Image Anal. 2017;36:61–78.

35. Tajbakhsh N, GURDUI SR, LIANG J. Automated polyp detection in colonoscopy videos using shape and context information. IEEE Trans Med Imaging. 2016;35(2):630–44.

36. Tajbakhsh N, GOLZAY M, LIANG J. Computer-aided pulmonary embolism detection using a novel vessel-aligned multi-planar image representation and convolutional neural networks. In: Navab N, Hornegger J, Wells W, Frangi A, editors. Medical Image Computing and Computer-Assisted Intervention – MICCAI 2015. Munich: Springer; 2015.

37. Balin SJ, Wetter DA, Andersen LK, Davis MD. Calcification of cutaneous disease occurring in association with autoimmune connective tissue disease: the Mayo Clinic experience with 78 patients, 1996-2009. Arch Dermatol. 2012;148(4):455–62.

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