

LETTER

Prediction of histology by B-mode and PD-mode ultrasound across different joint locations and diseases

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To cite: Micheroli R, Pauli C, Bürki K, *et al.* Prediction of histology by B-mode and PD-mode ultrasound across different joint locations and diseases. *RMD Open* 2022;**8**:e002439. doi:10.1136/rmdopen-2022-002439

► Additional supplemental material is published online only. To view, please visit the journal online (<http://dx.doi.org/10.1136/rmdopen-2022-002439>).

Accepted 27 June 2022



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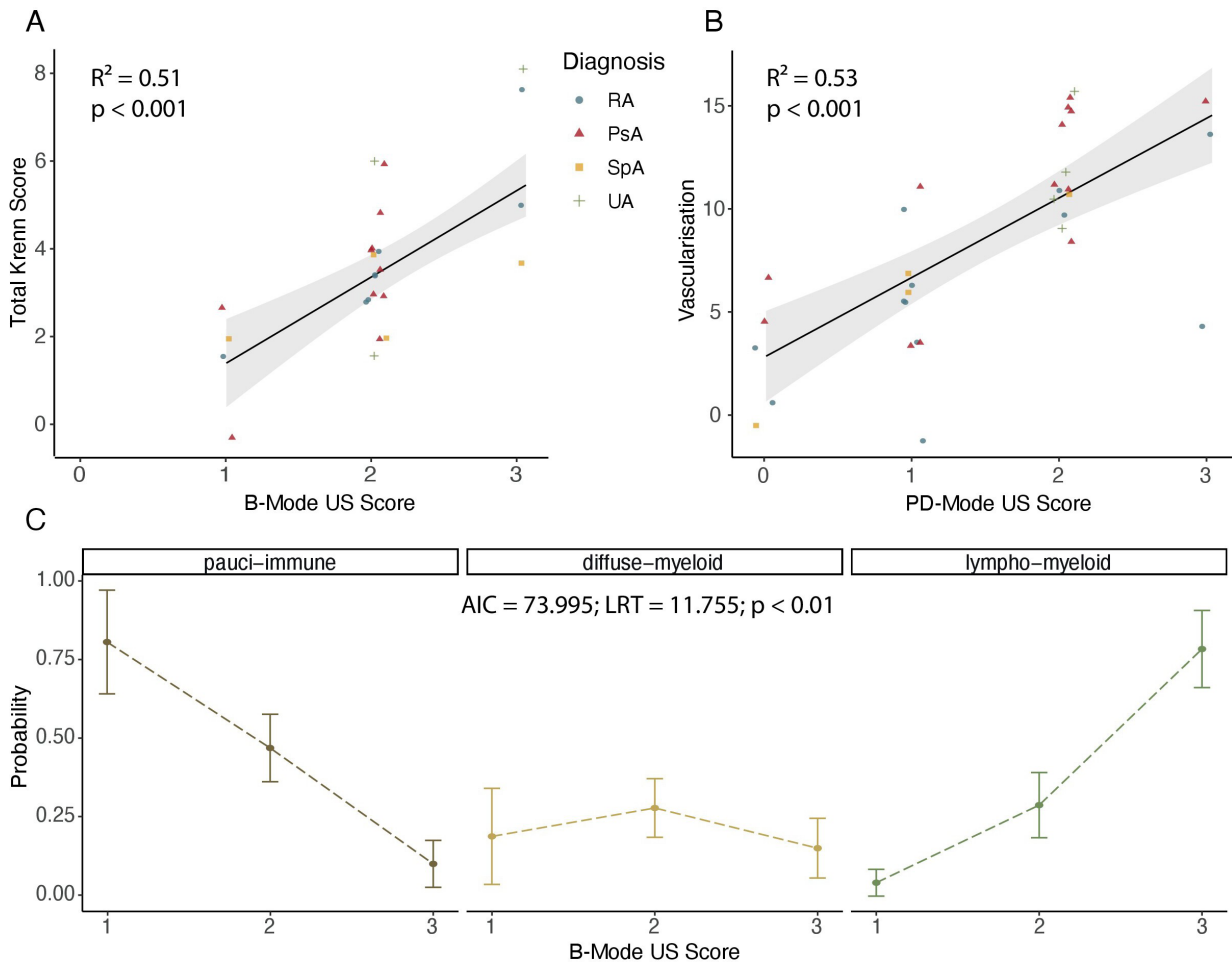
A few studies have investigated the predictive value of ultrasound (US) regarding specific histological findings and no study so far assessed the respective impact of different joint locations and diseases.^{1–3} Thus, the aim of this study was to investigate whether US—using grey scale (B)-mode and power Doppler (PD)-mode—can predict synovial characteristics in patients with specific inflammatory rheumatic diseases across different joint locations.

Ultrasound guided synovial biopsy (USGB) and histological examination were performed according to EULAR guidelines^{4 5} including questionnaires on tolerability and side effects before and after the procedure; ethical approval as well as informed consent was obtained (KEK2021-00092). Histology included quality control, Krenn Score,⁶ synovial pathotype according to Humby *et al*,⁷ and vascularisation based on the mean number of ERG positive blood vessels over five randomly selected visual fields ($\times 20$ magnification). B-mode and PD-mode US were scored semiquantitatively according to OMERACT definitions.⁸ Patients fulfilled classification criteria for the respective diseases. Statistical analyses included univariable and multivariable linear and multinomial logistic regression analysis (goodness-of-fit by likelihood ratio χ^2 test) and were performed with R V.4.1.3.

A total of 33 joints with available US scores and representative histology were assessed from 33 patients with the following diseases: rheumatoid arthritis (RA; n=12), psoriatic arthritis (PsA; n=13), spondyloarthritis (n=4) and undifferentiated arthritis (UA; n=4). USGB was very well tolerated with no side effects, and all patients would repeat the procedure, if necessary.

Participants' ages ranged from 20 to 79 years (mean 51, SD=13.75) and 64% of patients were female. The following joint locations were included: 10 wrists, 3 metacarpophalangeal, 18 knee joints, 1 sternoclavicular and 1 metatarsophalangeal joint. Median B-mode and PD-mode score were 2. A significant positive relationship between B-mode US and total Krenn score (**figure 1A**), and between PD-mode US and vascularisation (**figure 1B**) was found in univariate linear regression analysis. This effect remained significant in the adjusted analysis, including age, sex (reference men), disease (reference RA) and location (reference knee) as covariables (online supplemental table S1). In addition, PsA was related to significantly higher vascularisation in the multivariable model (estimate 3.27; 95% CI 0.19 to 6.36; p=0.039), a finding previously described.⁹ A multinomial logistic regression model using B-mode US as independent variable showed a high likelihood ratio for prediction of pathotype (LRT 11.76, p<0.01). Low B-mode US scores increased the probability of having a pauci-immune pathotype, while patients with high B-mode US scores had a high probability of having a lymphomyeloid pathotype (**figure 1C**). Adding total Krenn Score as additional variable to the multinomial model, B-mode US likelihood ratio diminished to 4.06 (p=0.13) suggesting the possible role of Krenn Score as mediator for the capability of B-mode US Score to predict the synovial pathotype.

In conclusion, this study suggests, that B-mode and PD-mode US scores according to definitions introduced for hand joints by OMERACT⁸ can predict the grade of histological synovitis and vascularisation independent of the disease analysed or of



joint location. Although B-mode showed a high likelihood ratio to predict the synovial pathotype, the identification of the diffuse-myeloid subtype seems to be challenging based on US characteristics alone. Additional clinical/imaging surrogate markers might, therefore, be needed.

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Contributors RM, CO and AC were involved in conception and design of the work. RM performed US examinations and joint biopsies, analysed the data and wrote the manuscript. CP did the histological analysis. PR performed US examinations and joint biopsies. KB organised and assisted all biopsies. OD, CO and AC contributed to the interpretation of the data and development of the manuscript. All authors critically revised the manuscript and approved the final version. CO and AC are joint last authors.

Funding The research leading to these results has received funding from Novartis Foundation for Biomedical Research, Stiftung Marie-Lou Ringgenberg, Stiftung für Rheumaforschung and the Swiss National Science Foundation (No. 320030_176061).

Disclaimer The funding sources had no role in the design and conduct of the study; collection, management, analysis, and interpretation of the data; preparation, review, or approval of the manuscript; or decision to submit the manuscript for publication.

Competing interests RM reports payment/honoraria from AbbVie, Eli Lilly, Gilead Sciences and Pfizer. CP, KB and PR have no CI to declare. OD reports consulting fees from Abbvie. CO reports payment/honoraria from CAS sex-specific and

gender-specific medicine, travel support from advances in targeted therapies. AC reports payment/honoraria from Abbvie, Merck Sharp & Dohme and Novartis.

Patient consent for publication Not applicable.

Ethics approval This study involves human participants and was approved by Local ethics commission of Zurich (KEK2021-00092). Participants gave informed consent to participate in the study before taking part.

Provenance and peer review Not commissioned; externally peer reviewed.

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