Use of Radiomics to Improve Diagnostic Performance of PI-RADS v2.1 in Prostate Cancer

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Objective: To investigate whether a radiomics model can help to improve the performance of PI-RADS v2.1 in prostate cancer (PCa).

Methods: This was a retrospective analysis of 203 patients with pathologically confirmed PCa or non-PCa between March 2015 and December 2016. Patients were divided into a training set (n = 141) and a validation set (n = 62). The radiomics model (Rad-score) was developed based on multi-parametric MRI including T2 weighted imaging (T2WI), diffusion weighted imaging (DWI), apparent diffusion coefficient (ADC) imaging, and dynamic contrast enhanced (DCE) imaging. The combined model involving Rad-score and PI-RADS was compared with PI-RADS for the diagnosis of PCa by using the receiver operating characteristic curve (ROC) analysis.

Results: A total of 112 (55.2%) patients had PCa, and 91 (44.8%) patients had benign lesions. For PCa versus non-PCa, the Rad-score had a significantly higher area under the ROC curve (AUC) [0.979 (95% CI, 0.940–0.996)] than PI-RADS [0.905 (0.844–0.948), P = 0.002] in the training set. However, the AUC between them was insignificant in the validation set [0.861 (0.749–0.936) vs. 0.845 (0.731–0.924), P = 0.825]. When Rad-score was added to PI-RADS, the performance of the PI-RADS was significantly improved for the PCa diagnosis (AUC = 0.989, P < 0.001 for the training set and AUC = 0.931, P = 0.038 for the validation set).

Conclusions: The radiomics based on multi-parametric MRI can help to improve the diagnostic performance of PI-RADS v2.1 in PCa.

Keywords: artificial intelligence, radiomics, prostate cancer, multi-parametric MRI, PI-RADS v2.1

INTRODUCTION

Prostate cancer (PCa) remains the most commonly diagnosed malignancy among men in the western world (1). The frequency of PCa in Asia has increased rapidly in years (2). Accurate detection and diagnosis of PCa are key factors to improve its therapeutic response and prognosis. Currently, magnetic resonance imaging (MRI) is generally considered the best modality for the detection and localization of PCa, and are thus becoming increasingly important (3). A recently developed multi-parametric (mp) MRI protocols including T2-weighted (T2W), diffusion-weighted
(DWI), and dynamic contrast-enhanced (DCE) imaging, appears to have good performance for PCa diagnosis, when associated with the Prostate Imaging Reporting and Data System (PI-RADS) (4).

In 2012, the initial version (v1) of PI-RADS was released to promote standardized MRI techniques and image interpretation. However, limitations of version 1 were soon evident. In 2015, version 2 was described to further improve reporting accuracy, and now has seen a broad uptake (4, 5). In 2019, PI-RADS version 2.1 was newly described, with several studies suggesting that version 2.1 could be preferable than version 2 for the evaluation of transition zone PCa (6–8). PI-RADS is now playing an increasingly prominent role in PCa diagnosis (9, 10). However, PI-RADS seems to have limitations of relatively low specificities, and inter-reader reproducibility. Thus a quantitative diagnostic method is needed to improve the performance of PI-RADS for the definite diagnosis of PCa (11, 12).

Radiomics can provide large-scale radiological image analysis by using a large number of quantitative features (13). Compared with genomics and proteomics, radiomics has the advantages of non-invasion assessments, comprehensive views of whole tumor and convenience in routine practice; thus, this technique has great potential for application in individualized diagnosis and treatment. Two studies (4, 14) have shown that radiomics can be used to detect PCa. However, it’s uncertain whether radiomics can add value to PI-RADS in the diagnosis of PCa. Therefore, this study aimed to determine whether radiomics of mpMRI can enhance the performance of PI-RADS v2.1 in PCa diagnosis.

MATERIALS AND METHODS

Patients

This study was approved by the local Institutional Review Boards (No. 2019-1209, Date: December 30, 2019) and the need for written informed consent was waived.

The institutional database of medical records was searched for suitable patients between March 2015 and December 2016. A total of 203 patients (mean age 66 years, age range 36–85 years) who met the following criteria were finally enrolled. The inclusion criteria: (1) Men with suspicious lesions on mpMRI; (2) These lesions were histologically confirmed by biopsy or radical prostatectomy; (3) no prior prostate endocrine therapy, biopsy, surgery, or radiation therapy before MRI examination. The exclusion criteria: (1) lesions with maximum transverse diameter ≤5 mm, which could hardly be delineated on MRI; (2) poor mpMRI quality. The patient recruitment pathway was shown in Figure 1.

The baseline characteristics, pathological data, and radiographic evaluation of each patient, including age, size, location, Gleason score, PI-RADS v2.1 score and prostate-specific antigen (PSA) were shown in Table 1. The patients were divided into two groups (the training and validation sets) at a ratio of 7:3 according to the scanning date.

MRI Examination

All MRI examinations were performed on the same 3.0 T MRI scanner (Skyra, Siemens Healthcare Sector, Germany) with a pelvic phased array coil. Scan sequences included T2WI in the axial and sagittal planes, DWI with b values of 0, 200, 400, and 1,000 s/mm², and DCE. ADC maps were calculated on a designated workstation. Supplementary Material summarizes the parameters of mpMRI sequences, including the type, repetition time/echo time (TR/TE), section thickness, field of view (FoV), and bandwidth.

Reference Standard for Pathology

All lesions were histopathologically proven based on biopsy (transrectal ultrasound [TRUS]-guided 12-core systematic biopsy) or surgical specimens (radical prostatectomy). Pathological confirmatory reports were acquired from medical records of the Department of Pathology.

PI-RADS Evaluation

Two experienced radiologists (more than 5 years of experience in the diagnosis of PCa) were assigned to review the mpMRI. The patient identification was removed from all images, and the readers were blinded to all clinicopathological information. The mpMRI including T2WI, DWI with corresponding ADC map, and DCE of the largest lesion in each patient was scored with a scale of 1–5 using PI-RADS v2.1. PI-RADS scores obtained by the two readers were assessed by a weighted Kappa statistics test to evaluate the inter-observer variability. Then any disagreement between the two readers was solved by discussion during the image interpretation.

Texture Feature Extraction and Model-Building

The images were normalized before feature calculation. In detail, each image was subtracted by the mean value and was divided by the standard deviation value. Then the image was multiplied by 100, and resampled to the same resolution.

Two radiologists drew volume of interest (VOI) independently on MR images of 30 patients to evaluate the stability of the features. Only the features with inter- and intra-class correlation coefficient (ICC) > 0.75 can be included in the following analysis. The entire VOI of the tumor were drawn on the base of radiologic-histologic correlation slice by slice (the radiologists were blinded to the histopathology results). For the patient with multiple lesions, only the dominant lesion (the largest lesion) was segmented.

Radiomic features of the lesions were extracted using PyRadiomics. Three types of features (first-order statistics, texture features, and shape features) for a total of 1,304 features were extracted from each sequence of mpMRI. To eliminate the differences in the value scales of radiomics features, all of the features were normalized before feature selection. Each feature was subtracted by the mean value of the training group and was divided by the standard deviation value. The same normalization method was applied to the validation set. Redundant features were removed by One-way analysis of variance (ANOVA). Then, the least
MRI examinations were performed on the same scanner between 2015 and 2016 (n = 282)

Patients included preliminarily (n = 219)

Finally included patients (n = 203)

PCa group (n = 112) Non-PCa group (n = 91)

Excluded:
- 25 treated with surgery or endocrine therapy before MRI
- 38 no pathological results
- 9 poor quality of mpMRI
- 7 small tumor volume

This table summarized the integrated PI-RADS scoring results for the two readers. The weighted Kappa between two readers was 0.722 (95% CI 0.651–0.794), which indicated strong consistency.

PCa, prostate cancer; PSA, prostate-specific antigen.

**TABLE 1** | Patient and tumor characteristics.

| Characteristics                  | PCa (n = 112) | Non-PCa (n = 91) | P value | Training cohort (n = 141) | Validation cohort (n = 62) | P value |
|----------------------------------|---------------|------------------|---------|---------------------------|---------------------------|---------|
| Age (y)                          | 66.0 ± 7.6    | 66.0 ± 10.3      | 0.970   | 66.5 ± 8.6                | 64.9 ± 9.42               | 0.245   |
| Size (long axis, mm)             | 2.22 ± 1.08   | 1.56 ± 0.88      | 0.001   | 1.86 ± 0.96               | 2.07 ± 1.21               | 0.194   |
| Location                         |               |                  |         |                           |                           |         |
| Transitional zone                | 51            | 56               | 0.023   | 72                        | 35                        | 0.479   |
| Peripheral zone                  | 61            | 35               |         | 69                        | 27                        |         |
| PSA (ng/ml)                      |               |                  |         |                           |                           |         |
| PSA ≤ 10                         | 15            | 41               | 0.000   |                           |                           | 0.749   |
| 10<PSA ≤ 20                      | 31            | 37               |         | 47                        | 21                        |         |
| PSA>20                           | 66            | 13               |         | 57                        | 22                        |         |
| Gleason score                    | 11/69/8/24    |                  |         | 8/50/5/15                 | 3/19/3/9                  | 0.797   |
| Extracapsular invasion on MRI (+/-)| 74/38        | 6/85             | 0.000   | 58/83                     | 22/40                     | 0.448   |
| PI-RADS v2.1 score               |               |                  |         |                           |                           |         |
| 1                                | 0             | 3                | 0.000   |                           |                           | 0.651   |
| 2                                | 0             | 26               |         |                           |                           |         |
| 3                                | 6             | 30               |         |                           |                           |         |
| 4                                | 32            | 25               |         |                           |                           |         |
| 5                                | 74            | 7                |         |                           |                           |         |

**FIGURE 1** | Flow chart of patients' recruitment pathway.
absolute shrinkage and selection operator (LASSO) regression method was applied to select the most distinguishable features.

Each clinical feature was assessed by univariate logistic regression. The features revealed as statistically significant with univariate logistic regression analysis were then analyzed with multivariate logistic regression analysis for model-building. A nomogram was generated for model visualization. Receiver operating characteristic (ROC) curve analyses were conducted to estimate the diagnostic performance of the models for the diagnosis of PCa.

Statistical Analysis
All statistical analyses were performed on R software, Statistic Package for Social Science version 21, Stata 15.0, and Medcalc 15.2.2. Differences in Table 1 were assessed by the chi-square test, the Mann-Whitney test, or t-test. The AUCs between different models were compared by DeLong’s test. The confidence level was set at $P < 0.05$.

RESULTS
A total of 203 patients were included in this study, in which 112 patients had PCa, and 91 patients had benign lesions [84 benign prostatic hyperplasia (BPH), and 7 high-grade prostatic intraepithelial neoplasia (HGPIN)]. For the lesion origin, 96 lesions originated from the peripheral zone, and 107 lesions were located in the transitional zone. The PCa had a larger size than non-PCa (long axis, $2.22 \pm 1.08$ mm vs. $1.56 \pm 0.88$ mm, $P = 0.001$). As shown in Table 1. As for the reference standard, 82 patients with benign lesions did not undergo radical prostatectomy, and the pathological results were determined by TRUS-guided biopsy.

Feature Selection and Model-Building
For the consistency test of VOIs, the number of features with ICC $> 0.75$ were 522 for DWI, 655 for ADC, 471 for DCE, and 266 for T2WI, as shown in Figure 2A. A total of 45 features (5, 19, 10, and 11 features were extracted from T2WI, DWI, ADC, and DCE images, respectively) were selected by LASSO method (Figure 2B). These features all had high ICC ($> 0.75$). The radiomics model (Rad-score) was comprised of these features in a formula shown in Supplementary Material. Rad-score had statistical difference between PCa and non-PCa groups (Rad-score = $0.85 \pm 0.29$ vs. $0.12 \pm 0.25$, $P < 0.001$). Then the combined models were built by combining PI-RADS (odds ratio [OR] = 6.4, $P = 0.001$) with Rad-score (OR = 14.5, $P < 0.001$) or PSA using multivariate logistic regression analysis.

A nomogram was generated for the combined model (PI-RADS + Rad-score) visualization (Figure 3A). To use the nomogram, find the point for each feature on the corresponding axis, add the points for all features, and draw a line from the total points axis to the risk axis to determine the risk of PCa. Higher total score was associated with greater risk of PCa. The model yielded satisfactory fit measurement based on the training set (Hosmer-Lemeshow test, $P = 0.943$). Moreover, there were also good calibration curves for the risk estimation (Figures 3B, C).

PCa vs. Non-PCa Classification
Rad-score had a significantly higher AUC [0.979 (95% CI, 0.940–0.996)] than PI-RADS [0.905 (0.844–0.948), $P = 0.002$] in the training set. However, the AUC between them was insignificant in the validation set [0.861 (0.749–0.936) vs. 0.845 (0.731–0.924),
When Rad-score was added to PI-RADS, the performance of PI-RADS was significantly improved for the PCa diagnosis (AUC = 0.989, \( P < 0.001 \)) for the training set and AUC = 0.931, \( P = 0.038 \) for the validation set). As shown in Table 2 and Figure 4.

In the case of clinical features, PSA, location, size, and extracapsular invasion were selected by univariate logistic regression analysis. However, only PSA was included in the combined model by multivariate logistic regression analysis. However, the addition of PSA to the combined model failed to show incremental diagnostic value (AUC = 0.990 vs. 0.989 for the training set; AUC = 0.937 vs. 0.931 for the validation set). As shown in Table 2 and Figure 4.

For differentiating the peripheral zone lesions, the AUC of the combined model was higher than that of PI-RADS (AUC = 0.995 vs. 0.891, \( P = 0.002 \)) for the training cohort; AUC = 0.941 vs. 0.753, \( P = 0.046 \) for the validation cohort; AUC = 0.987 vs. 0.860, \( P < 0.001 \) in the whole cohort). For differentiating lesions in the transitional zone, the AUC of the combined model was higher than that of PI-RADS (AUC = 0.981 vs. 0.909, \( P = 0.007 \) for the

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**Figure 3** (A) The radiomics nomogram was developed in the training cohort, with the Rad-score and PI-RADS incorporated. (B) Calibration curve of the nomogram in the training set. (C) Calibration curve of the nomogram in the validation set. Calibration curves depict the calibration of the model in terms of the agreement between the predicted risks of PCa (the x-axis) and observed outcomes of PCa (the y-axis). The blue solid line represents the performance of the nomogram (Note: a closer fit to the diagonal dotted line represents a better prediction).
training cohort; AUC = 0.926 vs. 0.861, \( P = 0.170 \) in the validation cohort; AUC = 0.960 vs. 0.894, \( P = 0.006 \) in the whole cohort).

**Classification of PI-RADS 3 Lesions**

Thirty-six patients in this study had PI-RADS 3 lesions on prostate MRI, in which 6 patients had PCa, and 30 patients had non-PCa. The Rad-score, and the combined model (Rad-score + PI-RADS) both had good diagnostic performance for the identification of PI-RADS 3 lesions (both AUC = 0.944), which was shown in Table 3 and Figure 5.

**DISCUSSION**

In this study, we proposed a combined model (PI-RADS combined with Rad-score) to significantly improve the diagnostic value of PI-RADS v2.1. By adding Rad-score to PI-RADS, the combined model outperformed PI-RADS in the diagnosis of PCa (AUC = 0.989 vs. 0.905, \( P < 0.001 \) for the training set; AUC = 0.931 vs. 0.845, \( P = 0.038 \) for the validation set). Thus adding quantitative Rad-score can benefit radiologists in the diagnosis of PCa.

PI-RADS v2 is designed to improve lesion detection, localization, characterization, and risk stratification in patients with suspected cancer (15). It is known that PI-RADS v2 generally benefits from its highly structured criteria, making relatively high diagnostic sensitivity in PCa diagnosis (14). However, the specificity and inter-reader reproducibility are moderate (16–18). To address these limitations of PI-RADS v2, an updated version (PI-RADS v2.1) was developed in 2019. One of the major modifications in version 2.1 is the diagnostic criteria for the transitional zone PCa of low T2WI scores. When comparing the performance between version 2 and 2.1 for characterization of suspected PCa, several studies suggested that AUC tended to be higher in version 2.1 than in version 2 without statistical significance (6–8). It must be noted that PI-RADS v2.1 still showed a high false positive rate (moderate specificity) for PCa diagnosis, similar to that with PI-RADS v2. Moreover, PI-RADS 3 lesions are frequently encountered (22–32%), and carry a moderate malignant potential (up to 20–30%), the stratification of these lesions is still challenging when using PI-RADS (19). Therefore, quantitative parameters, such as radiomics, may help to prevent misdiagnoses and improve performance of PI-RADS v2.1.

| Model                                               | The training cohort | \( P \) value | The validation cohort | \( P \) value |
|-----------------------------------------------------|---------------------|---------------|-----------------------|---------------|
|                                                     | AUC   | SEN  | SPE   | AUC   | SEN  | SPE   | AUC   | SEN  | SPE   |
| PI-RADS*                                           | 0.905 | 96.2%| 63.5% | 0.845 | 91.2%| 67.9% |
| Rad-score                                           | 0.979 | 89.7%| 96.2% | 0.002 | 82.4%| 82.1% | 0.825 |
| PI-RADS + Rad-score                                 | 0.989 | 92.3%| 98.4% | \(<0.001\) | 79.4%| 96.4% | 0.038 |
| PI-RADS + Rad-score + PSA                           | 0.990 | 92.3%| 98.4% | \(<0.001\) | 79.4%| 96.4% | 0.026 |

*When PI-RADS v2.1 score of the lesion \( \geq 4 \), this lesion was diagnosed as PCa.

\( P \) value, compared with PI-RADS.

AUC, Area under the curve; SEN, sensitivity; SPE, specificity.

PI-RADS v2 is designed to improve lesion detection, localization, characterization, and risk stratification in patients with suspected cancer (15). It is known that PI-RADS v2 generally benefits from its highly structured criteria, making relatively high diagnostic sensitivity in PCa diagnosis (14). However, the specificity and inter-reader reproducibility are moderate (16–18). To address these limitations of PI-RADS v2, an updated version (PI-RADS v2.1) was developed in 2019. One of the major modifications in version 2.1 is the diagnostic criteria for the transitional zone PCa of low T2WI scores. When comparing the performance between version 2 and 2.1 for characterization of suspected PCa, several studies suggested that AUC tended to be higher in version 2.1 than in version 2 without statistical significance (6–8). It must be noted that PI-RADS v2.1 still showed a high false positive rate (moderate specificity) for PCa diagnosis, similar to that with PI-RADS v2. Moreover, PI-RADS 3 lesions are frequently encountered (22–32%), and carry a moderate malignant potential (up to 20–30%), the stratification of these lesions is still challenging when using PI-RADS (19). Therefore, quantitative parameters, such as radiomics, may help to prevent misdiagnoses and improve performance of PI-RADS v2.1.
Compared with qualitative or subjective explanation of radiological images, radiomics permits high-throughput extraction of quantitative features to evaluate the degree of intratumor heterogeneity (20). In recent years, radiomics analysis has appeared as a potent tool for constructing decision-support models. A number of studies have used radiomics analysis to automate PCa diagnosis and risk stratification (21, 22). While few studies (4, 14) focused on the comparison of the diagnostic value between radiomics and PI-RADS. Our results suggested that the AUC of radiomics was higher than that of PI-RADS in the training set, which was consistent with previous studies. However, the difference of AUC was insignificant in the validation set, which was not entirely consistent with the previous studies. This finding showed that radiomics might not replace PI-RADS currently. The building methods between our study and those of previous studies were all machine learning, and all these studies lacked external validation. However, there were still some differences that need to be explained. Firstly, the version of PI-RADS in our study was 2.1, which was different from version 2 in previous studies (4, 14). Secondly, Chen et al. (14) only used T2WI and ADC images, in which lack of enhanced images reduced the effectiveness of mpMRI and radiomics. Finally, prior studies (4, 14) had the smaller sample sizes than ours, especially Wang et al. (14).

In our study, extracapsular invasion, location of lesions, and tumor diameter did not present enough predictive power for the differentiation of benign and malignant lesions. Thus, we integrated PSA into the combined model (Rad-score + PI-RADS). However, adding PSA failed to show incremental diagnostic value. This might be because the AUC of the combined model was high enough. In our work, when Rad-score were added, the diagnostic performance of PI-RADS was prominently improved: the specificity increased from 63.5 to 98.4% in the training set, and from 67.9 to 96.4% in the validation set. Adding Rad-score to PI-RADS might overcome the challenge of moderate specificity of PI-RADS. For the individual zone-based analysis, the combined model outperformed PI-RADS in the training, validation, and whole sets for differentiating lesions in the peripheral zone. However, the combined model failed to show significantly higher diagnostic performance in differentiating transitional zone lesions for the validation cohort (P = 0.170). We speculated that this might be related to the small sample size of transitional zone lesions in the validation set.

For PI-RADS 3 lesions classification, our exploratory results may provide preliminary evidence to justify the use of radiomics in this field. In clinical practice of the future, the validation of radiomics is important for the challenging PI-RADS 3 lesions, including biopsy or short-term follow-up in these lesions with high risk of PCa indicated by radiomics.

Our study had several limitations. First, due to its retrospective design, there might be selection bias between PCa and non-PCa groups, and the high b-value images (b value \( \geq 1,400 \text{ s/mm}^2 \)) failed to be obtained. Second, prospective and external validation was not performed. Third, all mpMRI images were obtained from a single institution. In the future, multicenter verification is needed to extend the versatility of the experimental results.

In conclusion, although the radiomics model cannot replace PI-RADS currently, adding radiomics to PI-RADS has the potential to improve the performance of the structured PI-RADS scheme by providing radiologists with quantitative and

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**TABLE 3** | ROC analyses of PI-RADS, Rad-score, and the combined models for identifying PI-RADS 3 lesions.

| Model                        | PI-RADS 3 lesions (n = 36) | \( P \) value |
|------------------------------|-----------------------------|---------------|
| AUC                          | SEN                         | SPE           |
| PI-RADS*                     | 0.500                       | 0             | 100%          | <0.001         |
| Rad-score                    | 0.944                       | 100%          | 80%           | <0.001         |
| PI-RADS + Rad-score          | 0.944                       | 100%          | 80%           | <0.001         |
| PI-RADS + Rad-score + PSA    | 0.911                       | 83.3%         | 90.0%         | <0.001         |

*When PI-RADS v2.1 score of the lesion \( \geq 4 \), this lesion was diagnosed as PCa.

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**FIGURE 5** | A 70-year-old man with a PI-RADS 3 lesion. (A) T2WI, heterogeneous signal intensity with obscured margins (arrow), which indicated 3 points according to PI-RADS v2.1; (B) ADC with 3 points, focal hypointense (discrete and different from the background); (C) DCE, contemporaneously with enhancement of adjacent normal prostatic tissues. According to (A–C), this lesion was classified as PI-RADS 3, indicating this lesion had an intermediate likelihood of clinically significant cancer. However, considering Rad-score (value = 0.97) and PI-RADS (score = 3), the risk of PCa was very high (the probability was about 85%) according to the nomogram. Finally, this lesion was proven to be PCa pathologically, with Gleason score = 3 + 4.
standardized criteria, thereby enabling us to more confidentially detect prostate cancer.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by West China Hospital of Sichuan University Biomedical Research Ethics Committee. Written informed consent for participation was not required for this study in accordance with the national legislation and the institutional requirements. Written informed consent was not obtained from the individual(s) for the publication of any potentially identifiable images or data included in this article.

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AUTHOR CONTRIBUTIONS

BS and LY conceived of the presented idea. ML and YY collected the data. ML, JX, and CH analyzed the data. ML drafted the manuscript. All authors reviewed the manuscript and BS made corrections to the manuscript. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fonc.2020.631831/full#supplementary-material
Conflict of Interest: Authors JX and CH were employed by the company Beijing Deepwise & League of PHD Technology Co., Ltd.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.