FULL PAPER

Diagnostic yield of CT-guided lung biopsies: how can we limit negative sampling?

1MARCELLO ANDREA TIPALDI, 1EDOARDO RONCONI, 2MILTIADIS E KROKIDIS, 1ALEKSEJS ZOLOVKINS, 1GIANLUIGI ORGERA, 1FLORINDO LAURINO, 1JULIA DAFFINA, 1DAMIANO CARUSO, 1ANDREA LAGHI and 1MICHELE ROSSI

1Department of Surgical Medical Sciences and Translational Medicine, Sapienza University of Rome - Sant’Andrea University Hospital, Rome, Italy
21st Department of Radiology, Aretaion University Hospital, Medical School, National and Kapodistrian University of Athens, Athens, Greece

Address correspondence to: Dr Marcello Andrea Tipaldi
E-mail: marcelloandrea.tipaldi@uniroma1.it; tipaldi.andrea@gmail.com

Objectives: To investigate whether lesion imaging features may condition the outcome of CT-guided lung biopsy (CTLB) and to develop a scoring system of biopsy outcome prediction.

Methods: This is a single center retrospective study on 319 CTLBs that were performed in 319 patients (167 males/152 females, mean age 68 ± 12.2). Uni- and multivariate analysis were performed aiming to assess the imaging features that are likely to be correlated to a negative biopsy outcome and patients were stratified in groups accordingly.

Results: Technical success was 100%. 78% of the biopsies (250/319) led to a concrete histology report (218 malignant/32 benign). The remaining lesions led to concrete histology at a second attempt that occurred on a later time. Multivariate analysis revealed increased risk of inconclusive result for nodules with low fludeoxyglucose uptake [odds ration (OR) = 2.64, 95% confidence interval (CI) 1.4–4.97; p = 0.003], for nodules with diameter smaller than 18 mm (OR = 2.03, 95% CI 1.14–3.62; p = 0.017) and for nodules that are located in one of the lung bases (OR = 1.96, 95% CI 1.06–3.62; p = 0.033). Three different groups of patients were identified accordingly with low (<30%), medium (30–50%) and high (>50%) probability of obtaining an inconclusive biopsy sample.

Conclusion: This study confirms that percutaneous CT-guided biopsy in nodules that are either small in diameter or present low positron emission tomography-fludeoxyglucose uptake or are in one of the lung bases may lead to inconclusive histology. This information should be factored when planning percutaneous biopsies of such nodules in terms of patient informed consent and biopsy strategy.

Advances in knowledge: Inconclusive histology after lung biopsy may be subject to factors irrelevant to technical success. Lung biopsy histology outcomes may be predicted and avoided after adequate planning.

INTRODUCTION

Percutaneous CT-guided lung biopsy (CTLB) is a first-line, minimal invasive, image-guided procedure implemented for the histologic characterization of focal lung lesions. Lung cancer screening and increased use of CT led to a raise in the detection rate of lung masses; most of such masses would require further characterization as further management is not possible without a histological confirmation of pathology. However, in some cases, CTLBs may be inconclusive. There are some arbitrary criteria on which lesions might lead to a non-conclusive result when biopsied; however, there is no clear consensus given the heterogeneity of evidence in the literature.

The aim of this study is to assess whether we may predict biopsy performance outcomes based on imaging features of the lesion. This information to our view is considered important in terms of following the appropriate biopsy strategy but also in terms of informed consent towards the patient that might need to undergo a repeated biopsy procedure.
Methods and materials

Patients
This is a single center retrospective study concerning the period between January 2014 and January 2019. Approval from the local ethics committee was obtained. We included in the study all the patients that underwent a CT-guided biopsy in our institution (Sant’Andrea University Hospital, La Sapienza, Rome, Italy) in the mentioned period and had in their records a fludeoxyglucose positron emission tomography (FDG PET) no longer than 6 months prior to the biopsy. Patients who underwent other procedures between the FDG PET and the biopsy were excluded. Out of the 1178 CTLBs that were performed in the mentioned period, 319 patients fulfilled the inclusion criteria and were included in the study.

Biopsy procedure
All biopsies were performed by four interventional radiologists with more than 5 years of experience on percutaneous CT-guided interventions; a 16 slice (Mx8000 IDT, Philips Healthcare, Cleveland, OH, EE. UU) or a 256 (Brilliance ICT 256, Philips Healthcare, Cleveland, OH, EE. UU) multislice spiral scanner was used for the procedures.

Patients were placed in supine, prone or lateral positions according to the most suitable approach to the lesion and a low-dose helical CT examination was performed in order to plan the access. Needle pathway, when possible, was delineated, avoiding interlobular fissures, visible bronchi, and relatively large vessels. Necrotic areas of the lesions (based on CT and FDG PET findings) were not targeted.

During needle advancement, images were acquired intermittently using CT fluoroscopy, which allowed single acquisitions to confirm and adjust the needle position. Full-Core Coaxial needle systems 18/20 Gauge were used in all cases. One to fourings) were not targeted.

A second unenhanced low-dose helical CT performed at the end of the intervention to detect possible early complications.

Data collection
Based on the obtained CT scan, lesion size, morphology and position including length of needle pathway from pleural surface to lesion were recorded. The latter was evaluated taking into account the optimal needle trajectory avoiding big size vessels, visible bronchi and interlobular fissures. A lesion was considered as located in one of the lung’s basal segments if a part of the diaphragm was included in the same CT frame as the lesion.

The standardized uptake value (SUV) was deducted from the PET report and an SUVmax equal or smaller to 2.5 was considered as the cut-off value for the purposes of this study, to consider a nodule as of low FDG PET uptake.

Data regarding each procedure have been extracted from the hospital patient records: total session duration, procedural time for biopsy and dose–length product (DLP) for each patient were collected. DLP was calculated considering the DLP of each scan and the number of scans performed in CT fluoroscopy.

Potential complications were assessed with chest plain films 3 h post–biopsy and when necessary, a CT scan.

Definition of outcomes
Technical success was defined as the evidence of the needle tip in the target lesion and the achievement of material susceptible of histological evaluation.

Histopathology reports were classified in three categories: positive for a specific malignant disease, positive for a specific benign disease and not adequate. The latter were deemed those, which resulted in necrosis, clots, normal parenchyma tissue, or those with not enough material for diagnosis.

Statistical analysis
Continuous variables were reported as mean (standard deviation, SD) or median (interquartile range, Q1–Q3). Distribution was checked for normality using the Shapiro–Wilks test. Categor- ical variables were presented as absolute number (n) and relative percentage frequency (%). To compare continuous variables between two groups, a two-tailed t-test was used when normally distributed; when the variables were not normally distributed, the non-parametric Mann–Whitney test was used. Pearson χ² was performed to test the association between categories for dichotomous variables. The continuous variables, maximum lesion diameter and distance from the pleural surface were dichotomized around an optimal cut-off identified on receiver operating characteristic curves applying Youden’s index (the maximum value of sensitivity + specificity).

We considered not adequate outcome as the response variable and performed uni- and multivariable analyses applying a logistic regression model with cluster robust estimate of standard error, considering the repetition of some patients. The multivariable logistic model was built considering as independent predictors those which resulted significant at a level of significance of 0.10.

The results were reported in terms of odds ratio (OR) and relative 95% confidence interval (95% CI). The predictive accuracy of the model was assessed through the c-index and its confidence interval (95% CI); performing the Hosmer–Lemeshow test assessed the goodness-of-fit of the model. The Bootstrap resampling method was chosen for internal validation of predictive models by selecting 300 repetitions. For each group of 500 bootstrap samples, the model was reassembled and tested against the observed sample in order to estimate predictive accuracy and distortion. As regards to calibration, the bootstrap was used to obtain correct estimates (corrected for overfitting) of the expected values compared to the observed ones represented in the calibration graph. The multivariable logistic model has been represented through a nomogram, a graphic representation that traces the effect of each variable to a scale ranging from 0 to 100, in order to add the scores obtained by the variables in the model, on the basis of the values observed, and to have a total risk score that corresponds to a greater or lesser probability of a
Table 1. Patient and nodule demographic features

| Population | N = 319 |
|------------|--------|
| **Age, years** | Mean ± SD: 68 ± 12.2 | Range: 32–88 |
| **Sex** | m: 185 | f: 134 |
| **Lesion diameter, mm** | Median (IQR): 23.25 (20.15) | Range: 5–120 |
| **Distance from the pleural surface, mm** | Median (IQR): 14 (29) | Range: 0–75 |
| **PET** | Negative, n: 70 | Positive, n: 249 |
| **Basal localization** | Yes, n: 75 | No, n: 244 |
| **Nodule aspect** | Solid, n: 263 | Subsolid, n: 40 | GG’s, n: 16 |

IQR, interquartile range; PET, positron emission tomography; SD, standard deviation.

Results of the statistical analyses are reported in Table 1. Technical success was 100%. A concrete histological outcome was obtained in 78% (n = 250) of the 319 cases. There were 218 (87%) malignant and 32 (13%) confirmed benign lesions. Malignant lesions were mainly adenocarcinomas in 54% of the cases (n = 118), squamous cell carcinomas in 16% (n = 35) and metastatic lesions in 12% (n = 26). The rest of malignant histology was distributed among lymphomas (n = 9), non-differentiated tumors (n = 9), small cell carcinoma (n = 10), carcinoid (n = 5), mesothelioma (n = 4) and sarcoma (n = 2). Benign diagnosis included chronic inflammation and organized pneumonia (n = 14), necrotizing and non-necrotizing granulomatosis (n = 6), reactive lymphoid hyperplasia (n = 4), hamartochondromas (n = 4), solitary fibrous pleural tumor (n = 2) and lesions positive for the DNA of Mycobacterium tuberculosis (n = 2). Non-conclusive histology occurred in the 22% (n = 71) of patients due to inadequate samples.

The mean session duration was 19 (±9 SD) min and the median duration of needle position and sampling was 8 min (interquartile range = 6–13). The median DLP value for the CT fluoroscopy time was 49.4 Gy x cm² (interquartile range = 33.8–75.4).

Complications were classified according to the CIRSE Guidelines. In our population, we observed Grade 1 complications such as pneumothorax not requiring intervention, limited parenchymal hemorrhage and transient hemoptysis, Grade 2 complications such as pneumothorax or iatrogenic bleeding requiring patient observation with overnight stay, and Grade 3 complications such as major pneumothorax treated via, chest tube placement.

Overall complications rate was 45%, however in 40% of the cases complication was only Grade 1 (in 23% small pneumothorax that did not require any further intervention or prolonged hospital stay and in 22% small alveolar hemorrhage). In 5% pneumothorax that required the placement of drainage tube and prolonged hospital stay occurred (Grade 2–3).

Nodule 18F-FDG-uptake and correlation with histology

The SUV max threshold of 2.5 was used as an arbitrary value to distinguish between PET “positive” (SUV max > 2.5- Group A) and PET “negative” (SUV max ≤ 2.5 - Group B) nodules. In total, 250 (78.4%) lesions were included in Group A and 69 (21.6%) in Group B.

Conclusion histology outcomes were significantly higher for Group A (82% vs 61%, p = 0.0003) and radiation exposure was significantly higher for Group B (median DLP 66.3 mGy x cm vs 48.8 mGy x cm, p = 0.0008).

Analysis of variables correlated with non-conclusive histology and predictive model

Lesions were analyzed in terms of maximum diameter and distance from the pleural surface. A cut-off value of 18 mm was considered (after dichotomization of values) for maximum lesion diameter and of 20 mm for distance from the pleural surface (Figure 1 and Table 2).

On the basis of the univariable analysis results, a multivariable logistic regression model was developed taking into account as independent variables the following: age at biopsy, FDG PET uptake, lesion localization in one of the lung bases, nodule diameter and distance from the pleural surface (Table 3). The results revealed a significantly higher adjusted probability of inconclusive sample for FDG PET negative nodules (OR = 2.64, 95% CI 1.4–4.97; p = 0.003), for nodules with diameter smaller than 18 mm (OR = 2.03, 95% CI 1.14–3.62; p = 0.017) and for nodules with localization in one of the two lung bases (OR = 1.96, 95% CI 1.06–3.62; p = 0.033). No significance was noted for the following parameters: (a) distance from the pleural surface smaller than 20 mm (1.65, 95% CI 0.9–3.03; p = 0.107) and (b) age at biopsy (OR = 0.98, 95% CI 0.97–1; p = 0.165).

Multivariable model was graphically depicted by a nomogram (Figure 2), obtaining the probability of failure occurring for a given individual. In the nomogram, each variable in the model was listed separately with a corresponding number of points assigned to a given magnitude of the variable. A total of 78 points were allocated to PET, 57 points to nodule diameter, 54 points to
basal localization, 40 points to pleural distance ≤20 mm, and an increasing number of points (0–100) the lower the patient's age. The cumulative point score for all the variables was matched to a probability of occurring failure. Three groups of low (<30%), medium (30–50%) and high (>50%) risk of failure patients, were finally detected with corresponding cut-off points of <142, 142–210 and >210. A total of 262 patients were allocated in the low-risk group (83%), 45 in the medium risk group (14%) and 12 in the high-risk group (3%). The model reveals that the parameter "low FDG PET uptake" appears high correlation with inconclusive biopsy results. In our population, it was present in 100% of the high-risk group, in 75% of the medium group and in only 5% of the low-risk group. The other two parameters that appear correlated to histological outcome are the lesions diameter (smaller than 18 mm) and the localization of the lesion in one of the two lung bases that were present in 25, 70, 87% and 17, 39,73% of the patients in the respective risk groups. As an example, we can observe how a patient 62 years old (33 pts), with a PET negative nodule (78 pts), basal located (54 pts) with a diameter ≤18.02 mm (57 pts) had a total score of 222, corresponding to the high-risk group (Figure 3).

**DISCUSSION**

Percutaneous CT-guided biopsy of lung lesions is a well-established minimal invasive percutaneous procedure for the characterization of focal lung lesions. Even though success is high, and most lesions may be easily targeted under CT guidance in most cases, histology outcome is not always conclusive. A potential strategy for this issue may be the use of a larger diameter biopsy system or the increase of the number of obtained cores, however, this might come with the additional cost of higher complication rate such as pneumothorax. It is therefore very useful to be knowledgeable on which lesions are more likely to lead to a non-conclusive result when biopsied.

Our study was successful in identifying those intrinsic features of lung nodules that are more likely correlated with a biopsy failure. Overall success rate of CTLB performed in our cohort are comparable to those previously reported in literature, especially considering the mean lesion diameter of 2.3 cm in our cohort, with some degree of variability between groups. In particular, the three features that resulted significant in our study were the following: basal localization, dimension of the lesion and the 18F-FDG uptake.

Diameter of the lesion was found to be a determining factor in our series, and this is in line with the previous report of Huang et al who found the diagnostic accuracy of CTLB to be significantly lower in nodules <15 mm, as in other older series. However, other authors reported rare difference in the diagnostic accuracy of CTLB for small nodules such as less than 20 mm and 8 mm, respectively. In our opinion, there are several possible explanations for these findings: type of needle used and biopsy technique, as previously suggested by Laurent et al, differences in population selection and the definition of “successful” procedures could be implicated.

| Cut Off | Sensitivity | Specificity | AUC | AUC 95% CI |
|---------|-------------|-------------|-----|------------|
| Diameter, mm | 18.02 | 49.30% | 68.15% | 58.70% | 52% | 65.3% |
| Distance from the pleural surface, mm | 20 | 70.42% | 41.94% | 56.2% | 50% | 52.3% |

AUC, area under the curve; CI, confidence interval; ROC, receiver operating characteristic.

More specifically, a cut-off of 18 mm was identified for maximum lesion diameter and of 20 mm for distance from the pleural surface.
Another factor that may condition the outcome may be the number of samples for each lesion. In our study, the number of samples was collected and analyzed preliminary, but we did not find statistical correlation with success. However, due to non-rigorous documentation in the reports it wasn’t possible to collect these data for all the procedures and we decided not to include it in the final research.

Basal localization of the nodules is a well-known limiting factor of percutaneous lung biopsies given the vicinity with diaphragm and the inevitable mobility of the lesions due to the respiratory motion. In our series, it resulted significant in the univariate analysis with a value of $p = 0.01$ and in the multivariable logistic regression with a value of $p = 0.033$. Equally, lesion dimensions less than 18 mm resulted significant in the univariate analysis with a value of $p = 0.007$ and in the multivariable logistic regression with a value of $p = 0.017$.

Low $^{18}$F-FDG uptake was the factor, which was most significantly linked with inconclusive outcome ($p<0.001$ in the univariate and $p = 0.003$ in the multivariate analysis). An explanation of this outcome may be the low cellular density present in some malignant lesions or the benign nature of the tissue, both factors correlated with a lower $^{18}$F-FDG uptake. Moreover, it’s well known that lung biopsies tend to perform worse in assessing benignity rather than malignancy.\(^8\)

The predictive model was successful in identifying three groups of patients with different probability of failure respectively low, medium and high risk. Other two features were included to build the model: the distance from the pleural access less the 20 mm and the per year increment in age at biopsy.
The low-risk group account mostly PET positive nodules greater than 18 mm for which biopsy appears to be linked to a very low risk of failure (<30%). In our population, this group counted 262 patients with a concrete histological diagnosis of 81%.

The high-risk group, on the other hand, are mostly patients with PET-negative basal nodules, smaller than 18 mm and present a very high risk of failure (>50%). In our population, this group counted 12 patients with a concrete histological diagnosis of 44%. We recognize that this group of patients is, usually, low populated (12 cases in our series) and represents exceptional and demanding request to our service to reach a certain diagnosis.

The group with medium failure risk (range between 30 and 50%) counted 45 patients in our series and presented a concrete histological diagnosis of 67%, which is quite low too (Figure 4). Even for these patients, we believe that CTLB should be carefully evaluated instead of other strategies. When applying the score to our population, we can classify our findings as follows.

![Figure 3](image1.png)

The high-risk class - preprocedural CT shows a 16 mm solid nodule with basal localization in a 62-year-old patient. (b) $^{18}$FDG PET-CT of the nodule (white arrow) shows poor metabolic activity (SUVmax 2.4), noticeable when compared to the liver (white arrowhead). According to our scoring system, total points allocated for this lesion were 222 (33 for the patient age; 54 for localization in the lung basis; 57 for small dimension <18 mm; 78 for PET negative result). (c) Biopsy of the nodule shows needle tip at the center of the target (red circle), yet collected material resulted not diagnostic due to scarcity of the material. $^{18}$FDG, 18-fluorodeoxyglucose; PET, positron emission tomography; SUV, standardized uptake value.

![Figure 4](image2.png)

The medium-risk class - image shows a 24 mm solid nodule with a smaller satellite nodule located in the lower right lobe in a young patient (46 years old). (b) $^{18}$FDG PET-CT of the nodule highlights an increased metabolic activity (SUVmax 6.8). Total points allocated to this lesion were 150 (40 + 54 + 56). (c) Biopsy of the nodule revealed the malignant nature of the lesion (adenocarcinoma). $^{18}$FDG, 18-fluorodeoxyglucose; PET, positron emission tomography; SUV, standardized uptake value.

![Table 4](image3.png)

Table 4. Risk score applied in our population

|                      | Low risk | Medium risk | High risk | $p$-value |
|----------------------|----------|-------------|-----------|-----------|
| Outcome failure (%)  | 19       | 33          | 56        | 0.005     |
| Procedure length (min) Mean ± SD | 19 ± 9   | 20 ± 8      | 28 ± 11   | 0.006     |
| Delivered dose (DLP) Mean ± SD | 56.8 ± 36.9 | 74.9 ± 48.6 | 106.8 ± 62.3 | <0.001 |
| Complication rate    | 43%      | 47%         | 67%       | 0.678     |

DLP, dose-length product; SD, standard deviation. N.S. = Not significant at level 0.05.
population we can observe how the failure rate increased proportionally in the three groups \((p = 0.005)\), as well as the procedure length and the delivered dose \((p = 0.006; p < 0.001)\) (Table 4).

Our study investigated which lesion imaging features condition the outcome of CTLB to minimize the inconclusive procedures. Ceci et al.\(^2\) recently reported how PET/CT-guided biopsy of lung lesions led to fewer inconclusive biopsies than CT-guided biopsy, confirming how the \(^{18}\)F-FDG uptake cover a key role in the potential histological characterization. Another recent report by Caruso et al.\(^3\) point out the ability of Radiomics CT-texture analysis in differentiating malignant from benign lung nodules with low FDG uptake, enhancing the importance of radiological heterogeneity beyond FDG PET/CT and lung biopsy. In this scenario, future studies are needed to assess if Radiomics features may give more information about the biopsy outcome prediction. The main limitation of our study is the lack of an external validation of the predictive model.

In conclusion, our study demonstrated how PET negative lung nodules are more difficult to characterize with percutaneous lung biopsy than PET-positive ones. Moreover, our predictive model has successfully identified two groups of patients with medium and high risk of biopsy failure where the procedure should be carefully evaluated, considering its invasive nature and the risk of longer procedures with higher rate of delivered radiation dose.

**FUNDING**

Open Access Funding provided by Universita degli Studi di Roma La Sapienza within the CRUI-CARE Agreement.

**REFERENCES**

1. Laurent F, Montaudon M, Latrabe V, Bégueret H. Percutaneous biopsy in lung cancer. *Eur J Radiol* 2003; 45: 60–8. doi: https://doi.org/10.1016/S0720-048X(02)00286-3
2. Yun S, Kang H, Park S, Kim BS, Park JG, Jung MJ. Diagnostic accuracy and complications of CT-guided core needle biopsy of solid and part-solid lesions. *Br J Radiol* 2018; 91: 20170946. doi: https://doi.org/10.1239/bjr.20170946
3. Anzidei M, Porfiri A, Andrani F, Di Martino M, Saba L, Catalano C, et al. Imaging-Guided chest biopsies: techniques and clinical results. *Insights Imaging* 2017; 8: 419–28. doi: https://doi.org/10.1007/s13244-017-0561-6
4. MacMahon H, Naishid DP, Goo JM, Lee KS, Leung ANC, Mayo JR, et al. Guidelines for management of incidental pulmonary nodules detected on CT images: from the Fleischer Society 2017. *Radiology* 2017; 284: 228–43. doi: https://doi.org/10.1148/radiol.2017161659
5. Aberle DR, Adams AM, Berg CD, Black WC, Clapp JD. Reduced lung-cancer mortality with low-dose computed tomographic screening. *N Engl J Med* 2011; 365: 395–409. doi: https://doi.org/10.1056/NEJMoai1102873
6. Bach PB, Mirkin IN, Oliver TK, Azzoli CG, Berry DA, Brawley OW, et al. Benefits and harms of CT screening for lung cancer: a systematic review. *JAMA* 2012; 307: 2418–29. doi: https://doi.org/10.1001/jama.2012.5521
7. McMahon PM, Kong CY, Weinstein MC, Tramontano AC, Cipriano LE, Johnson BE, et al. Adopting helical CT screening for lung cancer: potential health consequences during a 15-year period. *Cancer* 2008; 113: 3440–9. doi: https://doi.org/10.1002/cncr.23962
8. Gould MK, Donington J, Lynch WR, Mazzone PJ, Midthun DE, Naishid DP et al. Evaluation of Individuals With Pulmonary Nodules: When Is It Lung Cancer? In: *ed Diagnosis and Management of Lung Cancer, 3rd ed.* American College of Chest Physicians Evidence-Based Clinical Practice Guidelines. 2013. e935–120143(Suppl):e935-e120S.Chest. doi: https://doi.org/10.1378/chest.12-2351
9. Callister ME, Baldwin DR, Akram AR, Barnard S, Cane P, Draffan J, et al. British thoracic Society pulmonary nodule Guideline development group; British thoracic Society standards of care Committee. British thoracic Society guidelines for the investigation and management of pulmonary nodules. *Thorax* 2015; 70(Suppl 2).
10. Schreiber G, McCrory DC. Performance characteristics of different modalities for diagnosis of suspected lung cancer: summary of published evidence. *Chest* 2003; 123(1 Suppl): 115S–28. doi: https://doi.org/10.1378/chest.123.1_suppl.115S
11. Duka E, Ierardi AM, Floridi C, Terrana A, Fontana F, Carrafiello G. The role of interventional oncology in the management of lung cancer. *Cardiovasc Intervent Radiol* 2017; 40: 153–65. doi: https://doi.org/10.1007/s00270-016-1495-y
12. Kohthary N, Lock L, Sze DY, Hofmann LV. Computed Tomography–Guided percutaneous needle biopsy of pulmonary nodules: impact of nodule size on diagnostic accuracy. *Clin Lung Cancer* 2009; 10: 360–3. doi: https://doi.org/10.1016/j.clc.2009.n.049
13. Gupta S, Wallace MJ, Cardella JE, Kundu S, Miller DL, Rose SC, et al. Quality improvement guidelines for percutaneous needle biopsy. *J Vasc Interv Radiol* 2010; 21: 969–75. doi: https://doi.org/10.1016/j.jvir.2010.01.011
14. Chang Y-Y, Chen C-K, Yeh Y-C, Wu M-H. Diagnostic feasibility and safety of CT-guided core biopsy for lung nodules less than or equal to 8 MM: a single-institution experience. *European Radiology* 2017; (2017).
15. Susam S, Çinkooğlu A, Ceylan KC, et al. Diagnostic success of transbronchial needle biopsy and PET-CT for 1 to 2 cm solid indeterminate pulmonary nodules. *Clin Lung Cancer* 2009; 10: 360–3.
16. Yamagami T, Yoshimatsu R, Miura H, Yamada K, Takahata A, Matsumoto T, et al. Diagnostic performance of percutaneous lung biopsy using automated biopsy needles under CT-fluoroscopic guidance for ground-glass opacity lesions. *Br J Radiol* 2013; 86: 20120447. doi: https://doi.org/10.1259/bjr.20120447
17. Filippadis DK, Binkert C, Pellerin O, Hoffmann RT, Krajina A, Pereira PL. CIRSE quality assurance document and standards for classification of complications: the CIRSE classification system. *Cardiovasc Intervent Radiol* 2017; 40: 1141–6. doi: https://doi.org/10.1007/s00270-017-1703-4
18. Heerink WJ, de Bock GH, de Jonge GJ, Groen HM, Vliegenthart R, Oudkerk M. Complication rates of CT-guided transbronchoscopic lung biopsy: meta-analysis. *Eur Radiol* 2017; 27: 138–48. doi: https://doi.org/10.1007/s00330-016-4357-8
19. Huo YR, Chan MV, Habib A-R, Lui I, Ridley L. Pneumothorax rates of CT-guided lung biopsy: a comprehensive systematic review and meta-analysis of risk factors. *Br J Radiol* 2020; 93: 20190866. doi: https://doi.org/10.1259/bjr.20190866
Diagnostic yield of CT-guided lung biopsies

20. Huang M-D, Weng H-H, Hsu S-L, Hsu L-S, Lin W-M, Chen C-W, et al. Accuracy and complications of CT-guided pulmonary core biopsy in small nodules: a single-center experience. *Cancer Imaging* 2019; 19: 5151. Published 2019 Jul 23. doi: https://doi.org/10.1186/s40644-019-0240-6

21. van Sonnenberg E, Casola G, Ho M, Neff CC, Varney RR, Wittich GR, et al. Difficult thoracic lesions: CT-guided biopsy experience in 150 cases. *Radiology* 1988; 167: 457–61. doi: https://doi.org/10.1148/radiology.167.2.3357956

22. Li H, Boiselle PM, Shepard JO, Trotman-Dickenson B, McLoud TC. Diagnostic accuracy and safety of CT-guided percutaneous needle aspiration biopsy of the lung: comparison of small and large pulmonary nodules. *AJR Am J Roentgenol* 1996; 167: 105–9. doi: https://doi.org/10.2214/ajr.167.1.8659351

23. Laurent F, Latrabe V, Vergier B, Montaudon M, Vernejoux JM, Dubrez J. CT-guided transthoracic needle biopsy of pulmonary nodules smaller than 20 mm: results with an automated 20-gauge coaxial cutting needle. *Clin Radiol* 2000; 55: 281–7. doi: https://doi.org/10.1053/crad.1999.0368

24. Chang Y-Y, Chen C-K, Yeh Y-C, Wu M-H. Diagnostic feasibility and safety of CT-guided core biopsy for lung nodules less than or equal to 8 MM: a single-institution experience. *Eur Radiol* 2018; 28: 796–806. doi: https://doi.org/10.1007/s00330-017-5027-1

25. Caruso D, Zerunian M, Daffina J, Polici M, Polidori T, Tipaldi MA, et al. Radiomics and functional imaging in lung cancer: the importance of radiological heterogeneity beyond FDG PET/CT and lung biopsy. *Eur J Radiol* 2021; 142: 109874. doi: https://doi.org/10.1016/j.ejrad.2021.109874

26. Juliano J, Cerci JJ, Bogoni M, Cerci RJ, et al. PET/CT-Guided biopsy of suspected lung lesions requires less Rebiopsy than CT-guided biopsy due to Inconclusive results. *J Nucl Med* 2021; 62: 1057–61.