Comparison of the whole slide imaging and conventional light microscopy in the grading of oral epithelial dysplasia: a multi-institutional study

Priscilla Barbosa Diniz 1, Marcondes Sena-Filho 1, Karen Mendes Graner 2, Bruno Augusto Linhares Almeida Mariz 1, Leonardo Amaral dos Reis 1, Oslei Paes de Almeida 1, Jacks Jorge 1

1 Department of Oral Diagnosis, Piracicaba Dental School, University of Campinas (Unicamp), Piracicaba, SP, Brazil
2 Botucatu Medical School, São Paulo State University (UNESP), Botucatu, Brazil

Correspondence:
Department of Oral Diagnosis, Piracicaba Dental School
University of Campinas (UNICAMP)
901, Limeira Av., Areião, Piracicaba, São Paulo, Brazil
jacks@fop.unicamp.br

Received: 02/04/2020
Accepted: 28/10/2020

Abstract
Background: Whole Slide Imaging (WSI) is an alternative method to light microscopy (LM). However, few studies have compared the diagnostic agreement between WSI and LM, especially to grade oral epithelial dysplasia (OED). The purpose of this study was to evaluate the variability in grading OED by the World Health Organization grading system, using WSI and conventional LM, and to investigate whether the access to clinical information, and psychologic or physical states of the pathologists could interfere with the diagnosis.

Material and Methods: eleven experienced pathologists from seven Brazilian universities independently evaluated twenty-five OED cases. The analyses were performed in duplicate for each method, with an interval of at least 30 days, and the time consumed in each analysis was measured. Physical and psychologic states were evaluated by blood pressure levels, heart rate and two questionnaires: State-Trait Anxiety Inventory and Perceived Stress Scale. Clinical information was provided after the second evaluation using WSI and the pathologist could change their diagnostic decision or not.

Results: LM showed a higher inter-examiner agreement (k=0.53) than WSI (k=0.45) and a smaller time consumed by the pathologists (mean of 65.53 seconds compared to 91.02 seconds in WSI). In the first analysis using conventional microscopy, there was a positive correlation between kappa values and anxiety (r=0.47, p=0.02), and stress (r=0.64, p<0.01), and an inverse correlation with heart rate (r=-0.48, p=0.02). In the digital analysis, there was also a positive correlation between kappa values and anxiety (r=0.75, p<0.001). After clinical information was given, there was a slight change in 11.3% of the cases, and a great discrepancy in 1.1% of the cases, mainly increasing the OED grade.

Conclusions: both microscopy systems had similar results, although LM had slightly higher kappa values, and WSI was more time consuming.

Key words: Pathology, microscopy, diagnosis, leukoplakia oral, anxiety.
Introduction
The term oral potentially malignant disorder (OPMD) describes clinically detected epithelial lesions that carry an increased risk of progression to cancer (1-3). Oral epithelial dysplasia (OED) is one of the most common types the OPMD and its grading and diagnosis are basically made by the histopathological investigation, and lesions with a high grade of dysplasia are considered to be at higher risk of malignant transformation (3,4). Performing a correct diagnosis is the most important role for pathologists, and the literature reports three main causes for diagnostic mistakes: systemic causes, cognitive mistakes and lack of knowledge (5). The emotional state can have a negative cognitive impact at the moment of decision as well as the subjective aspect of the lesion, which is a reality in OED (6-8). In this context, access to clinical information, lesion site, and clinical images are imperative in the diagnostic process (9). The use of different methods of microscopic visualization can also influence lesion grading or even the final diagnosis (10,11).

Light microscopy (LM) is the classic method used by pathologists to achieve an accurate diagnosis. However, new methods have been developed in the last decades (12), helping pathologists to capture, visualize, analyze, store and share slides electronically, which are increasingly and easily available in most pathology centers (13-15). Whole Slide Imaging (WSI) has been used in Pathology for teaching and diagnostic purposes, including pathology conferences, consultations, reviews, slide panels, and more recently, for the initial diagnosis and archiving (14,16,17).

Considering all the advantages of WSI and the currently easier access to this technology, some studies have tried to validate its use in histopathology settings, but limited studies have compared the diagnostic agreement between WSI and traditional LM (14). Hence, our study aimed to evaluate the intra observer variability in grading OED by the World Health Organization (WHO) grading system, and the effects that pathologists' physical and psychological states have in the diagnostic performance.

Material and Methods
- Sample selection
Cases diagnosed between 2000 and 2015 as hyperkeratosis and acanthosis, epithelial dysplasia (mild, moderate or severe) or in situ carcinoma were selected from the Oral Pathology department of the Piracicaba Dental School, University of Campinas. Only cases with complete clinical data (age, sex, location, time of evolution, diagnostic hypothesis and clinical photograph) were selected by two pathologists that did not participate in the subsequent analysis. Inclusion criteria included: the diagnostic agreement between both pathologists, no technique errors, tissue integrity, and adequate coloration. Twenty-five cases, including 5 cases of hyperkeratosis and acanthosis, 16 cases of OED (6 mild, 6 moderate and 4 severe) and 4 cases of in situ carcinomas attending these criteria were selected and classified according to the WHO grading system.
- WSI imaging
Slides stained with hematoxylin and eosin (HE) were coded and scanned using the slide scanner Aperio ScanScope CS (Aperio Technologies, Vista, CA, USA). All images were checked for focus, brightness and contrast balance. All WSI analyses were performed using the software ImageScope version 11.2.0 Aperio®, in the same computer (Ultrabook Asus S46C, Intel® Core™ i5-33175 CPU @ 1.70GHz, 8 GB RAM). In the LM analysis, the pathologists used their usual microscope.
- Microscopic analysis
Thirteen pathologists were invited to participate in this study, and eleven voluntarily accepted. Six meetings were scheduled with each pathologist, two for methodological instructions and four meetings for actual study. Each slide was diagnosed twice using WSI or LM. All pathologists filled a form informing their age, time working as an oral pathologist, institution, and level of education (graduation, post-graduation, international experience). Before analyzing the study cases, ten control cases (including mucocele, fibrous hyperplasia, and squamous cells carcinoma) were selected to familiarize the pathologists with the research method. Initially, the study cases were analyzed by conventional LM by each pathologist. After 30 days, the pathologists analyzed the same cases digitally. After a minimum of 30 days interval, the second course of evaluations was performed to compare the intra observer agreement. The time spent for each analysis was measured. Pathologists and researchers were blinded to the slides and images identification. The slides were randomly identified by codes and the display order was changed for each course of the evaluation.
To assess the influence of clinical information on OED grading, clinical photos and clinical information were provided after the pathologists completed the second course of WSI. The slides were evaluated again and a final diagnosis was established for each case.
- Stress and anxiety analyses
Pathologists were evaluated for the presence and severity of stress and anxiety symptoms. A complete version of the Perceived Stress Scale (PSS), in the version translated to Portuguese by Luft et al., [2007] (18), was used to evaluate the symptoms of perceived stress in the month before the analysis. Anxiety was determined using the State-Trait Anxiety Inventory (STAI), a scale with 40 items, equally divided into two subscales (19). The first subscale evaluates the anxiety state (STAI-S) and expresses the feeling in the present moment. The
second subscale is the trait (STAI-T) and evaluates stable aspects of anxiety tendency, such as general states of calmness and confidence (20).

STAI-T was measured only in the first meeting with the pathologists. STAI-S and PSS were measured in all meetings before the analysis of the slides.

- Physical analysis
After the pathologists answered the questionnaires, blood pressure and heart rate were measured using a digital upper arm blood pressure monitor (Omron HEM-7113). This measurement was performed in all meetings, right before and right after the analysis of the slides. Normal values of arterial pressure and heart rate were considered normal (111/75 mmHg) and 73 bpm.

- Statistical analysis
Data were analyzed using the software SAS version 9.3 (SAS Institute, Cary, NC). Weighted Cohen’s Kappa was used to measure the intra-observer agreement, and it was interpreted as follows: Kappa < 0.10 (no agreement); 0.11 to 0.40 (weak); 0.41 to 0.60 (regular); 0.61 to 0.80 (moderate); 0.81 to 0.99 substantial (great) and Kappa =1.00 (perfect agreement). Multiple comparisons and the Tukey-Kramer test, Wilcoxon test, dependent T-test (for dependent data) and Pearson’s test were used in this study for association with the other variables. The statistical significance was set at p < 0.05.

Results
- Participants characteristics
All participants were professors of Pathology in Brazilian Universities and were experienced in the histopathological routine. They were mostly males (54.4%), with a mean age of 46 years old (SD±10.77) and had an average of 19.8 years (SD±11.9) of working time. One pathologist had a medical degree and the other 10 were dentists; 27.3% of all participants were specialists in Pathology, almost all had MSc degrees (90.9%) and all had PhD diplomas. Also, 54.5% had international experience and 63.6% had always worked in the same laboratory.

- Intraobserver Agreement
The most frequent diagnosis of the study was moderate dysplasia (27.8%), followed by severe (20.9%) and mild dysplasia (20.6%), hyperkeratosis and acanthosis (16.5%), and carcinoma in situ (14.2%) (Fig. 1). Weighted Cohen’s Kappa showed a higher agreement rate in LM (k = 0.53, CI95% 0.46-0.60) than in WSI (k = 0.45, CI95% 0.38-0.51), representing a regular agreement for both methods. There was a variation of the diagnosis among pathologists by both methods, and this variation was statistically significant in both methods, but LM varied less (Chi-Square test p = 0.009) than WSI (Chi-Square test p = 0.0008).

- Time for diagnosis
The time for performing the diagnosis was evaluated for WSI and LM. OED; either mild, moderate or severe; consumed more time for diagnosis than other lesions. LM consumed on average 65.5 second for each case and was faster than WSI, which took a mean of 91.0 seconds (p < 0.0001, Tukey-Kramer test). Moreover, the time for the diagnosis of each slide varied between the phases of the study. We found that in the first phase 84.5 seconds were spent in each case, while in the second phase the necessary time decreased to 72.1 seconds (p=0.002, Tukey-Kramer test).

- Effect of clinical information
To evaluate the importance of clinical information in the diagnosis, the pathologists first observed each slide without the clinical information and established their diagnosis. Then, this information was provided and the pathologists could change the diagnosis if necessary. Correlating the diagnosis in these two moments, there was a slight change in 11.3% of the cases, and a great discrepancy in 1.1% of the cases after receiving the clinical information. In most cases where the clinical information influenced the diagnosis, the pathologists tended to increase the grade of dysplasia, since the mean difference between the diagnosis was -0.101 (p<0.01, Wilcoxon test).

Fig. 1: Frequency of diagnoses rendered using light microscopy and whole slide imaging.
Grading oral epithelial dysplasia using light microscopy or WSI

- Psychological characteristics
  We observed that the pathologists were slightly anxious, as showed in the results of STAI-T, with a median score of 29.36 (SD±3.668) (20). When STAI-S was applied after each evaluation, an increase in the anxiety was observed, characterizing moderate anxiety (33.91, SD±6.5115) (Fig. 2).
  The mean value of stress among the pathologists during this study was 19.47 (SD±6.65), characterizing mild stress. Stress levels were similar during all the study phases. There was only one outlier (Fig. 2).
  When correlating the physical and psychological evaluations with the diagnosis agreement (kappa values) using Pearson's test, only in the first stage there was a positive correlation between kappa values and anxiety ($r=0.47, p=0.02$), and stress ($r=0.64, p<0.01$), and an inverse correlation with heart rate ($r=-0.48, p=0.02$). In the digital analysis, there was also a positive correlation between kappa values and anxiety ($r=0.75, p<0.001$).

Discussion

The current study was conducted to analyze the variability on grading OED by the WHO grading system, using LM and WSI. Some studies compared WSI and LM using tissues from different anatomic sites, such as the gastrointestinal tract (11,12), breast (21), prostate (22), and skin (23). However, this is the first multi-institutional study comparing these two methods for the histopathologic grading of OED. Additionally, according to our results, LM still remains the main tool to evaluate and grade OED, presenting higher intraobserver agreement, and being less time consuming than WSI.

Cell morphology and tissue architecture are the main characteristics of grading OED in the diagnostic routine (24). Aspects such as nuclear atypia and mitotic figures are used to determine the degree of dysplasia (25). The WSI method may result in difficulty identifying these features due to poor image resolution at high magnification, although we did not find that image quality was a major factor in the diagnostic discrepancy. Also, the lack of experience of pathologists using the WSI system is considered a factor for the discrepancies in the diagnoses between WSI and LM (26).

On the other hand, the variability in the diagnosis of OED was already expected due to the subjectivity of each pathologist’s analysis, regardless the use of different technologies (4,25). In our study, both WSI and LM methods showed similar results regarding the final diagnosis. The concordance among the pathologists in our study ranged from $k=0.53$ in LM to $k=0.45$ in WSI and was similar to previous studies using LM (24,25).

The WSI method was more time-consuming than LM in other studies (11,14,27), corroborating with our findings. This can be explained by lack of experience in the WSI method, and by the learning curve of new technologies (21). This limitation may discourage pathologists from adopting the digital platform in routine laboratory practice (21). We also observed that the first phase was more time consuming than the second, which corroborates the founds of Gui et al. (11), suggesting that the diagnosis of epithelial dysplasia using WSI may become more rapid with experience.

Generally, pathologists do not change their OED diagnosis even when they receive the patient’s clinical information. There are few studies about the influence of clinical information in OED (9). Abbey et al. demonstrated that absence of clinical information did not interfere with the pathologists’ diagnosis of OED. The authors reported that the results could be more accurate, but it would not change their conclusions (9).

The diagnostic routine involves experience, skills, fatigue and competency, factors that can influence the decision of the pathologist (6). For decisions with more than one potential interpretation, more anxious people
tend to behave more negatively than less anxious individuals with potential consequences for microscopic interpretation (7). We observed that in the first LM evaluation there was an increase in stress and anxiety after the diagnosis, which may have occurred due to tension or discomfort with a new and unknown situation (28). However, in the following evaluations, the pathologists became more confident with the methodology of the study. S-Anxiety was greater than T-Anxiety in all the stages of the study, which may be the result of an increase in attention levels (7).

WSI has been increasingly used for education, remote consultation, second opinions, and archiving (15,17,22). Despite the increased use of WSI in Europe and North America, there is a lack of available evidence to validate the use of WSI in routine primary diagnoses. The barriers to its implementation include cost-effectiveness, lack of evidence validating the diagnostic agreement and low acceptability among pathologists (27). Few institutions and laboratories have access and use this method in Brazil (29), although this method presents several advantages such as the off-site access to the digital slides, a quick and easy way to consult with other pathologists, to avoid lost or damaged slides, and enabling better ergonomics (30).

Some pathologists may find it more difficult than others to use WSI systems. We consider that our results show that pathologists can reliably grade OED using WSI, at similar levels to LM.

This is the first study to compare WSI with LM in grading OED. Our results indicate that both systems have similar results, although LM had slightly higher kappa values and was less time consuming. Conversely, more training and access to this new method could change these results. Further studies and more training are necessary to assess differences between WSI and LM in grading OED, and in other groups of oral lesions.

References

1. Warnakulasuriya S, Ariyawardana A. Malignant transformation of oral leukoplakia: A systematic review of observational studies. J Oral Pathol Med. 2016;45:155-66.
2. Shearston K, Fateh B, Tai S, Hove D, Farah CS. Malignant transformation rate of oral leukoplakia in an Australian population. J Oral Pathol Med. 2019;48:530-7.
3. Iosea O, Sollecito TP, Alawi F, Weinstein GS, Newman JG, De Virgilio A, et al. Potentially malignant disorders of the oral cavity and oral dysplasia: A systematic review and meta-analysis of malignant transformation rate by subtype. Head Neck. 2020;42:539-55.
4. Speight PM, Abram TJ, Floriano PN, James R, Vick J, Thornhill MH, et al. Interobserver agreement in dysplasia grading: Toward an enhanced gold standard for clinical pathology trials. Oral Surg Oral Med Oral Pathol Oral Radiol. 2015;120:474-82.
5. Ford J, Pambrou C. Exit competencies in pathology and laboratory medicine for graduating medical students. The Canadian approach. Hum Pathol. 2015;46:637-42.
6. Nag R, Kumar Das R. Analysis of images for detection of oral epithelial dysplasia: A review. Oral Oncol. 2018;78:8-15.
7. Hartley CA, Phelps E.A. Anxiety and Decision-Making Catherine. Biol Psychiatry. 2012;72:2.
8. Brennan M, Migliorati CA, Lockhart PB, Wray D, Al-Hashimi I, Axell T, et al. Management of oral epithelial dysplasia: a review. Oral Surgery, Oral Med Oral Pathol Oral Radiol Endodontology. 2007;103 Suppl:S19.e1-12.
9. Abbey LM, Kaugars GE, Gunsolse JC, Burns JC, Page DG, Svirsky JA, et al. The effect of clinical information on the histopathologic diagnosis of oral epithelial dysplasia. Oral Surg Oral Med Oral Pathol Oral Radiol Endod. 1998;85:74-7.
10. Al-Janabi S, Huisman A, Vink A, Leguit RJ, Offerhaus GJA, Ten Kate FJW, et al. Whole slide images for primary diagnostics of gastrointestinal tract pathology: A feasibility study. Hum Pathol. 2012;43:702-7.
11. Gui D, Naini B, Gerney G, Dry S, Cortina G, Hart S, et al. Diagnosis of dysplasia in upper gastrointestinal tract biopsies through digital microscopy. J Pathol Inform. 2012;3:27.
12. Al-Janabi S, Huisman A, Van Diest PJ. Digital pathology: Current status and future perspectives. Histopathology. 2012;61:1-9.
13. Ghanavi F, Evans A, Madabhushi A, Feldman M. Digital Imaging in Pathology: Whole-Slide Imaging and Beyond. Annu Rev Pathol Mech Dis. 2013;8:331-59.
14. Williams BJ, DaCosta P, Goacher E, Treanor D. A systematic analysis of discordant diagnoses in digital pathology compared with light microscopy. Arch Pathol Lab Med. 2017;141:1712-8.
15. Bertram CA, Klopfeisch R. The Pathologist 2.0: An Update on Digital Pathology in Veterinary Medicine. Vet Pathol. 2017;54:756-66.
16. Fonseca FP, Santos-Silva AR, Lopes MA, de Almeida OP, Vargas PA. Transition from glass to digital slide microscopy in the teaching of oral pathology in a Brazilian dental school. Med Oral Patol Oral Cir Bucal. 2015;20:e17-22.
17. Brierley DJ, Speight PM, Hunter KD, FartHING P. Using virtual microscopy to deliver an integrated oral pathology course for undergraduate dental students. Br Dent J. 2017;223:115-20.
18. Luft CDB, Sanches SdO, Mazo GZ AA. Brazilian version of the Perceived Stress Scale: translation and validation for the elderly. Rev. Saúde Pública. 2007;41:606-15.
19. Reidy J. Trait anxiety, trait depression, worry, and memory. Behav Res Ther. 2004;42:937-48.
20. Julian LJ. Measures of anxiety: State-Trait Anxiety Inventory (STAI), Beck Anxiety Inventory (BAI), and Hospital Anxiety and Depression Scale-Axiety (HADS-A). Arthritis Care Res. 2011;63:S467-72.
21. Krishnamurthy S, Mathews K, McClure S, Murray M, Gilmore M, Albarracin C, et al. Multi-institutional comparison of Whole slide digital imaging and optical microscopy for interpretation of hematoxylin-eosin-stained breast tissue sections. Arch Pathol Lab Med. 2013;137:1733-9.
22. Camparo P, Egevad L, AlbaGA F, Berney DM, Boccon-Gibod L, CompérAT E, et al. Utility of whole slide imaging and virtual microscopy in prostate pathology. Apnns. 2012;120:298-304.
23. Mooney E, KempF W, Jemec GBE, Koch L, Hood A. Diagnostic accuracy in virtual dermatopathology. J Cutan Pathol. 2012;39:758-61.
24. Manchanda A, Shetty DC. Reproducibility of grading systems in oral epithelial dysplasia. Med Oral Patol Oral Cir Bucal. 2012;17:e935-42.
25. Krishnan L, Karpagaselvi K, KumaRswamy J, Sudheendra US, Santosh K V, Patil A. Inter- and intra-observer variability in three grading systems for oral epithelial dysplasia. J Oral Maxillofac Pathol. 2016;20:261-8.
26. Pantanowitz L, Sinard JH, Henricks WH, Fatherree LA, Carter AB, Contis L, et al. Validating whole slide imaging for diagnostic purposes in Pathology: Guideline from the College of American pathologists Pathology and Laboratory Quality Center. Arch Pathol Lab Med. 2013;137:1710-22.
27. Goacher E, Randall R, Williams B TD. The diagnostic concordance of whole slide imaging and light microscopy: a systematic review. Arch Pathol Lab Med. 2016;141:151-61.
28. Racic M, Todorovic R, Ivkovic N, Masic S, Joksimovic B, Kulic M. Self-Perceived Stress in Relation to Anxiety, Depression and...
Health-related Quality of Life among Health Professions Students: A Cross-sectional Study from Bosnia and Herzegovina. Zdr Varst. 2017;56:251-9.

29. Pereira E Silva A, Nicolau Campos SM, Guimarães IM, Agate Platais Brasil Teixeira G. Comparison between digital and optical microscopy: Analysis in a mouse gut inflammation model. Biomed Reports. 2017;7:247-50.

30. Pantanowitz L, Evans A, Pfeifer J, Collins L, Valenstein P, Kaplan K, et al. Review of the current state of whole slide imaging in pathology. J Pathol Inform. 2011;2:36.

Acknowledgements
The authors are grateful for the pathologists who participated in this study: Fábio Daumas Nunes, Suzana Cantanhede Orsini Machado de Sousa, Andresa Borges Soares, Bruno Augusto Benevenuto de Andrade, Mário Romañach, Fábio Ramoa Pires, João Luiz de Miranda, Nádia Lages Lima, Ricardo Alves de Mesquita and Maria Cassia Ferreira de Aguiar. This study would not be possible without their time and effort.

Funding
Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES), finance code 001, and São Paulo Research Foundation (FAPESP) - grant #2018/24715-2.

Conflict of interest
None of the authors have any relevant financial relationship(s) with a commercial interest. This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

Ethics
The Ethics Committee in Research of the Piracicaba Dental School, University of Campinas approved the study (protocol no. 40348514.0.0000.5418).