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International Guidelines for Veterinary Tumor Pathology: A Call to Action

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1 International Guidelines for Veterinary Tumor Pathology: A Call to Action

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67 Introduction

68 Reproducibility is the foundation of experimental science but irreproducibility of published oncological studies is a crisis in human oncology and certainly also a problem 69 in veterinary oncology. In many instances the lack of reproducibility stems from 70 71 inadequate description of published methods. (Begley and Ellis 2012, Oransky Ivan 72 2017, Stark 2018, Wen, Wang et al. 2018) Efforts to address this crisis have been implemented in human medicine, including attempted reproduction of published studies 73 74 and formulation of checklists for use by journal editors and reviewers to ensure inclusion and transparency of detailed methods and materials in publications.(Oransky Ivan 2017, 75 76 Editorial 2018, Schott, Tatiersky et al. 2018, Stark 2018, Wen, Wang et al. 2018) Less 77 than 10% of observational studies are able to be replicated and incredibly, less than 78 20% of preclinical trials can be replicated. (Begley and Ellis 2012) If that is the state of 79 oncology studies in human medicine, how do we compare in veterinary oncology? How 80 far have we advanced in the last 40 years? Attempts to validate existing studies and or grading schemes are almost nonexistent. Grading schemes and the methods employed 81 82 require appropriate validation before they should be adopted and used to provide prognoses or direct clinical therapy. Yet our philosophy seems to be that once a system 83 84 or method is created, it is put in use and remains in use regardless of whether the system has been validated or not. We do not know how that system or method will 85 perform when different pathologists use it and when it is applied to new patients. 86 Consensus statements that support use of studies are not validation. Authors should 87 feel complimented when colleagues attempt to reproduce their methods and study 88 designs. Results will not replicate exactly, but our methods must. Validation of new 89 grading systems is impossible if the original methods cannot be duplicated by other 90 investigators. 91

92 **Reproducibility**

One of the major reasons that published studies are not able to be reproduced is the lack of sufficient details of the methods used to assess basic histological parameters including mitotic figure (MF) recognition, mitotic count (MC), lymphovascular invasion, tumor necrosis and margin evaluation.(Meuten, Munday et al. 2018, Schott, Tatiersky et

al. 2018) Currently, the assessment of these parameters requires pathologists to make 97 subjective evaluations without clearly defined methods. Due to the inconsistency of 98 these qualitative evaluations, there is weak or poor concordance between pathologists. 99 This may result in negativity of the parameters or rejection of the grading system. The 100 materials and methods section of manuscripts should contain descriptions of each 101 102 method in sufficient detail to allow others to reproduce the study and validate the results. Citing that the methods described in a prior published study were followed is 103 acceptable protocol, providing that any modifications used are described in detail. 104 Failure of studies to be replicated can be due to poorly described methods, not following 105 methods, and other confounders. Interobserver variation between pathologists reported 106 in manuscripts is often ascribed to a method that is inadequate or too subjective. 107 108 However, investigators may not have considered that the original methods were flawed or inadequately described, such that pathologists could not consistently follow the 109 method. Stating that MF were counted in 10 consecutive high-power fields (hpf) at 400X 110 is insufficient detail for others to reproduce the method, compare results and validate 111 112 the data. Since the area within one hpf can vary by greater than 200% because of different microscope configurations, then of course there will be interobserver variation 113 114 in MC if the microscopes used by study pathologists were not configured in the same way. Not only MC, but all parameters that were enumerated histologically (e.g. 115 116 pleomorphism) with a microscope or with whole slide images (WSIs) have the potential for error and variability of results if the area enumerated is not defined in mm². Until 117 methods are detailed such that others can reproduce them, we will have inconsistent 118 and contradictory data in the literature. Even with standardized methods, there will be 119 120 variability that needs to be reported and practical considerations that cannot be standardized. 121

122

123 Standardizing methodology is one step toward consistent results, but it does not 124 guarantee consistency and certainly not usefulness. The methods must be followed, 125 done carefully, using the same materials (e.g. antibody), and methods applied to 126 reference populations and diseased groups with known and well defined outcomes. 127 Accredited clinical pathology labs use standardized methods, calibration of instruments

and quality control measures to quantitate substances and report the reproducibility, 128 sensitivity, specificity, validation, and reliability, such as positive and negative predictive 129 values of test results. Similar principles need to be applied to anatomic pathology. 130 Quantitation of morphologic structures by semi- or fully automated means will need to 131 be validated and applied with similar rigid standards. (Boyce, Dorph-Petersen et al. 132 2010) When the methods are reproducible, they can be applied to cases with known 133 outcomes. It would be helpful if new methods to quantitate structures were compared to 134 the existing more subjective means to enumerate structures in HE stained slides.(Puri, 135 Hoover et al. 2019, Bertram, Aubreville et al. 2020) Then colleagues can compare 136 results to determine if they wish to adopt the new method. Technology will continue to 137 spur development of new methods that can be applied to diagnostic cases. Many 138 139 owners will pay for new techniques at any cost, but other owners will decline based on practical considerations such as cost, age of pet, or emotional value of the pet to their 140 141 family. How to balance best care with practicality of animal ownership is not simple. Researchers can help address this by comparing new methodologies with those that 142 143 can be performed without additional costs, specialized equipment, or expertise. Development of interlaboratory proficiency programs to promote standardization of 144 145 tumor grading system results and performance of ancillary testing, such as immunohistochemistry, is sorely needed. Although new methods may initially be 146 147 restricted to the institutions in which they were developed, standardization and proficiency are critical as these techniques are validated in other laboratories and 148 become routinely used for tumor diagnosis. Centers that develop novel tests (e.g. 149 computational pathology or CPATH), artificial intelligence, molecular, genetic) should 150 151 have a goal that the methods can be applied uniformly and are described in sufficient detail that other labs can perform and validate the tests. Newly developed, specialized 152 assays should be compared to current methodology and to patient outcomes to assess 153 their utility and ideally seek FDA approval. (Boyce, Dorph-Petersen et al. 2010, Puri, 154 Hoover et al. 2019, Bertram, Aubreville et al. 2020) 155

156 **Outcome assessment**

In addition to a lack of standardized assessment of histological criteria, 157 reproducibility in animal studies is also limited by a lack of standardized guidelines for 158 159 outcome assessments of animal patients.(Meuten, Munday et al. 2018, Schott, Tatiersky et al. 2018) Euthanasia unrelated to tumor progression appears to be a 160 significant confounder. Reported patient survival times are impacted by euthanasia 161 162 which may be elected due to personal decisions, varying judgements regarding quality of life, owner income or other factors which do not reflect tumor behavior. Furthermore, 163 the Start time (T=0) needs to be clearly and consistently defined in presented survival 164 analyses.(Nguyen) How often has an assigned grade or reporting that a tumor is in a 165 lymph node resulted in euthanasia and the patient may have lived significantly longer? 166 Survival time statistics in veterinary oncology are influenced by many factors outside of 167 168 tumor and host biology. Metastasis needs to be subdivided into confirmed or suspected. Evidence for metastases determined by imaging should be labelled as suspected or 169 170 *metastases as determined by imaging* when reported in journals. Histopathology is required to confirm that metastases are present and are of the same tumor type. 171 172 Multiple aggressive tumors can occur in the same patient and are well recognized in breeds such as Golden retriever, Rottweiler and Bernese mountain dogs.(Cullen and 173 174 Breen 2016) Oncology studies no longer routinely include results of autopsy, the perceived value of which seems to have hit a nadir. Owner/client permission to perform 175 176 autopsies should be pursued with sympathy and empathy but as vigorously as other tests. Autopsy findings greatly increase the objectivity of results such as metastases 177 178 and recurrence and therefore confidence of study results. Veterinary oncology studies need institutions and labs to pool their resources so that large numbers of cases can be 179 180 collected. If results of these studies are correlated with accurate patient outcomes, the 181 archived materials are a precious resource. The materials from these studies (slides, blocks, images, statistical data) could be shared with others such that new methods can 182 be applied to case series with known outcomes. This was done by Bergin et al. (Bergin, 183 Smedley et al. 2011) in a study of canine oral and lip melanocytic neoplasms. In 184 185 addition to a set of oral/lip melanocytic neoplasms from the authors' own diagnostic laboratory, this study used archived blocks from two previous studies(Spangler and 186 Kass 2006, Esplin 2008) in order to validate the histologic parameters described in 187

those studies (nuclear atypia, mitotic count, and pigmentation) using the same methods 188 and to compare them to a new parameter, Ki67 index. This can serve as an example of 189 190 the value in validating prior reports, which adds confidence to conclusions and provides a new method. Archived images of tumors could also be used to test inter-pathologist 191 variation on diagnoses, MC, necrosis, and other basic parameters from laboratories 192 193 worldwide. Outcome assessment determines if a test predicts treatment or prognosis and may differ between tertiary and primary care patients. Standardization of outcome 194 assessment data is as critical as the standardization of techniques involved in tumor 195 assessment. These two components of oncology must be linked as it is useless to 196 create a new tumor grading system without knowledge of patient outcomes 197

198

Appendices and Protocols

In 2011 Veterinary Pathology published a series of recommendations and 199 200 reviews about tumors in animals and how they should be evaluated. The manuscripts in that issue are excellent. They exceed the goals of the present manuscript but like "all" 201 veterinary pathology publications there is no provision to update information, which is 202 now a decade old. Much of the information consisted of literature reviews, and 203 descriptions of the multiple methods to perform a parameter, without prioritizing or 204 choosing one. The present manuscript aspires to be a continuum of the information 205 206 published in 2011 but with a focus on establishing standardized histopathology methods 207 to evaluate tumors. These methods are guidelines that will help accrue similar data such that studies can be cross compared and validated. A website will be established to 208 publish guidelines for standard methods of tumor evaluations with the purpose of 209 advancing veterinary pathology and oncology. This will require modifying the contents 210 211 when publications have substantial data driven results that warrant updating these 212 guidelines. These changes will be dated, and references cited. The present system of waiting for publication of a book or a fascicle is outdated. Updates are also needed as 213 214 errors are possible (authors are humans) in the present appendices and protocols, and it is possible that some important references were missed. The authors hope that 215 216 readers will bring such errors to our attention by contacting one or more of the communication authors. Unlike an error or omission in a manuscript or book that 217

remains in print, and results in our names indelibly associated with the words *retraction* or *correction*, the website can be quickly and easily updated. Journals and books will remain vital to our professions as they provide the means to publish peer reviewed research and to describe in detail an entire topic. The website will attempt to remain focused, and current, more of a CliffNotes' version of a topic designed to aid pathologists, editors and researchers in the standard parameters used to evaluate tumors and checklists of information that should be gathered about specific tumor types.

225 Appendices are guidelines to be used for identification of MF, perform MC, assess lymphovascular invasion (LVI), margin evaluation, percent tumor necrosis, 226 227 CPATH, lymph nodes and outcome assessments. These parameters have not been standardized for animal tumors. The methods are detailed for MF, MC, LVI and margin 228 229 assessments while others are newly developed methods (CPATH) or need clarification. 230 Tumor necrosis is used in grading systems for some tumors, yet the method to 231 determine percent necrosis in tumors from pets has never been described or not in sufficient detail such that others can reproduce the method (see Appendix 4). At the end 232 233 of each appendix is a section titled "Future Considerations", which provides a list of possible ways to improve that method. *Protocols* are designed to gather complete data 234 235 sets for the evaluation of commonly graded canine neoplasms. Protocols are provided for Soft Tissue Tumors/Soft Tissue Sarcomas (STT/STS), and are in process for canine 236 mast cell tumor, and canine melanoma (cutaneous and oral). Protocols for other tumors 237 can be developed and are needed, including mammary, splenic, osteosarcoma, 238 hematopoietic and lymphoid tumors as well as cytologic protocols. If we do not 239 standardize the methods used to identify tumors, we will continue to have conflicting 240 data in the literature. Protocols and Appendices can be used as guides for reviewers 241 242 and editors of manuscripts to ensure all required data was included and standard methods were followed. Journals serve as a gatekeeper for scientifically sound data, 243 and they should also not refrain from publishing negative results. Investigators can use 244 protocols as a checklist to ensure complete data sets are included for study participants. 245 The protocols are modeled after the College of American Pathologists with an emphasis 246 on gathering uniform data on specific tumor types. What are the consequences of not 247 following an appendix or protocol? Nothing, no accreditation or certification or plaques 248

of accomplishment will be awarded or rescinded. The methods described herein are 249 intended to be "best practices" that will add consistency and reproducibility to our 250 251 methods with an eye to our clients: clinicians, oncologists, patients, and the public. Appendices and Protocols extend beyond "best practices" as they provide brief 252 literature reviews, areas of weaknesses and list suggested fields of investigation for 253 254 future studies to improve a method. The guidelines described are based upon review of literature and authors' expertise, and are intended to bring consistency and 255 reproducibility to the evaluation of tumors in animals. These guidelines have not been 256 certified, accredited or reviewed by any standards-creating body and represent the 257 authors' own interpretation and application of the data reviewed. Application of these 258 guidelines may vary with different laboratories and personnel, and each pathologist 259 260 should consider whether these guidelines are appropriate based on the equipment, tissues or other materials available. Whether a governing body will aid in further 261 development in updating these guidelines will depend upon the success of the website 262 and how widely it is used. 263

264 **Future Collaboration**

The website being constructed will address some of these needs, but additional 265 personnel will be needed to maintain the site, develop different protocols, generate new 266 267 data, and validate studies. The initiative of a website with living appendices and tumor 268 protocols will be successful if others use this information in their diagnostic, research, and publication efforts and if the appendices and protocols are updated in a timely 269 manner as new information becomes available. A key benefit of standardization of 270 tumor evaluation is the ability to evaluate data accrued from studies of many 271 272 investigators at various institutions world-wide. This will permit analysis of larger data 273 sets and increase the statistical power of the observations. The eventual goal would be to develop veterinary pathology industry standards with international input and 274 275 acceptance. The goal is to accrue data on the important parameters that should be evaluated for a specific tumor type so that, over time, large data sets with comparable 276 277 information about specific tumor types can be evaluated to provide accurate prognostic information that improves patient care. This will take multi-institutional participation and 278

specialists from different disciplines. The driving force will likely come from younger 279 generations. Future appendices might include molecular profiles, genetic tests, and 280 281 checklists for surgical pathology reports. Protocols are unlimited, think of a tumor, write a protocol using these as templates. Edits and updates are encouraged: contact the 282 communication authors of an appendix or protocol. Submission of additional tumor 283 protocols is welcomed and can be accomplished by contacting the administrators of the 284 website. Confirming the need for standardized parameters to evaluate animal tumors 285 met with near unanimity. Agreement for the guidelines of each parameter is not always 286 unanimous. To compare data between labs, and ultimately improve patient care, we 287 need to apply the same methods to basic parameters used to evaluate tumors. Using 288 unstandardized methods that can cause variation in results is not scientifically sound. 289 290 Drawing conclusions for clinical cases based on methods that are not standardized is misleading. 291

292 Completed Appendices and Protocols are in the supplemental section of this 293 manuscript and they will be posted on the website and updated as needed 294 (www.vetcancerguidelinesandprotocols.cldavis.org). The following are excerpts and 295 summaries of each appendix or protocol, not the completed documents. Readers 296 interested in a parameter should read the details in completed documents provided in 297 the supplemental section of this manuscript and on the website.

298 SUMMARY

299 The goal of this project is to help advance veterinary oncology and pathology by promoting standardization of tumor assessment and patient outcomes. Guidelines are 300 proposed to increase the uniformity and consistency of methods used to evaluate 301 tumors along with suggestions for future consideration to help improve their 302 discrimination and utility. Scientific journals, editors and reviewers can ensure progress 303 in the goals of tumor assessment standardization and study reproducibility by 304 establishing certain requirements of manuscripts being reviewed. Oncology studies 305 306 which include histopathologic and gross features of tumors should have a pathologist as a co-author and journals should require this. Data obtained from record review without 307 knowledge of the diagnostic or grading criteria limits conclusions and confidence in the 308

study. Review of gross description and histologic slides or images by an authoring 309 pathologist or multiple pathologists is needed to ensure accuracy and uniformity of the 310 pathology data and that current methods and terminology are used. The appendices are 311 designed to help accomplish this. Certain appendices are completed: MF recognition, 312 MC, necrosis, LVI, margin assessment and synoptic reporting, while others are in 313 progress. The key steps to performing each method are condensed into checklists 314 within the appropriate appendix. These checklists should integrate well with synoptic 315 reporting (see Appendix 8). There are also discussion and notes to clarify certain steps. 316 The checklists for margin evaluation are subdivided by responsible persons, the list that 317 a pathologist should report are short and practical. LVI can be evaluated in HE sections 318 and methods to confirm and differentiate LVI from pseudo-vascular invasion (see 319 320 Appendix 3). Future investigations need to determine the importance of identifying if the tumor thrombus is in a lymphatic or blood vessel, and if the distinction has practical 321 322 importance it will need to be determined how capable pathologists are of distinguishing each type of vessel with HE stained sections. Some authors would like to see necrosis 323 324 abandoned as a parameter but that will require additional investigations. Suggestions to improve how necrosis is determined are provided in Appendix 4. CPATH will aid new 325 326 investigations and synoptic reporting will provide a means to summarize and readily retrieve information. Outcome assessments are central to improvement of prognostic 327 328 parameters but are under the umbrella of oncologists. However, histopathology is needed to confirm it is the same tumor in a recurrence or metastasis. 329

330 Until there are data driven results that can be standardized and proven prognostically useful, tumor assessment will need to include a wide range of parameters. Some 331 practices, such as reporting margins of benign tumors or mitotic counts in tumors in 332 which significance is not established will be left to the discretion of the pathologist and 333 clinician. Clinicians faced with decisions on patient therapy rely extensively on 334 pathologists' assessments. The prognostic significance of various factors changes over 335 time necessitating clarity in communication of pathological findings, giving clinicians the 336 information needed. The website is a window for clinicians to see pathologists' 337 338 perspective of tumor assessment. Fascial planes to the surgeon are not the same as to 339 the pathologist, a high power field is not a standard unit of area, if surgical margins are

not inked by the clinician, there is no accuracy to HTFD and there are other examples to
illustrate our different perspectives. We need to do our best for the clients, owner, and
pet, but we also need to explain and defend our discipline. The latter will be easier if
veterinarians entering our profession understand our roles, and the limitations of our
techniques.

345 The appendices and protocols require updating and renewal to be useful documents. Pathologists, oncologists and other scientists are encouraged to submit suggestions 346 347 and supporting data to enable thoughtful revision. Tumor types and behavior may differ in various geographic sites and we encourage communication from all points of the 348 349 globe to enhance our overall understanding of tumor behavior. Protocols are needed for additional tumor types and appendices should be developed for other parameters such 350 351 as cytological assessments to recognize and grade specific tumor types, cellular and nuclear pleomorphism and proliferative indices. Research needs to clarify which 352 technique and modifications enhance diagnostic and prognostic accuracy and if they 353 can be practically applied to diagnostic cases, and subsequently validated with robust 354 data. As in most research endeavors, new technology should be directed to answer 355 356 specific problems and not end up as a new method in search of a question to answer.

Prospective studies that follow rigorous guidelines are the standard we should strive for 357 358 and which will help guide the way forward. (Webster, Dennis et al. 2011) We also propose a platform from which new data can be gathered and integrated into an 359 360 ongoing approach to evaluate the practicality and utility of current, as well as newer 361 methods of tumor evaluation. Publishers can aid this project by providing permission for 362 authors to copy sections of manuscripts they authored without forcing them to rewrite their own sentences to avoid plagiarism. How long will it take to accomplish all of this is 363 364 unknown, but we need to continue and expand upon what our colleagues started in 2011. 365

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409

410 Mitotic count (MC) and Histologic Morphology of Mitotic Figures (MF) (See 411 Appendix 1 and Appendix 2)

MC will remain an important parameter in the evaluation of tumors as it is easy to 412 accomplish, incurs no additional costs, is predictive of tumor proliferation and is part of 413 414 multiple grading schemes that help predict tumor behavior. However, certain components essential for performing reproducible MC must be defined including the 415 region of the tumor where MC should be performed (ie "hot spot" or areas of highest 416 mitotic density within a tumor)(van Diest, Baak et al. 1992, Baak, , Meyer, Cosatto et al. 417 2009, Al-Janabi, van Slooten et al. 2013, , Veta, Van Diest et al. 2015, Meuten, Moore 418 et al. 2016, ,) and the amount of tumor area in which MF will be counted, expressed in 419 standard units of measure (ie mm²).(Meuten, Moore et al. 2016) Although enumerating 420 MF has long been a mainstay of tumor assessment, until recently there has been no 421 standardization of any element of this parameter in veterinary pathology. Perhaps, the 422 assumptions were that we were all counting the correct structures with the same 423 method, that these methods matched published manuscripts and therefore there was no 424 need to standardize the technique. Unfortunately, some of those assumptions are false. 425 Performing the MC is considered laborious but subjective with inter-pathologist 426 427 variation.(, Tsuda, Akiyama et al. 2000, Meyer, Alvarez et al. 2005, Veta, Van Diest et 428 al. 2015, Bertram, Gurtner et al. 2018,) Possible causes include counting differently sized areas, poorly defined methods, not following methods, counting too rapidly, 429 counting ambiguous structures, experience level, tumor mitotic heterogeneity, inability 430 to find hot spots, quality of sections (fixation, artifacts) and quality of images. 431

To achieve accurate and consistent counts the MC must be performed carefully following standardized procedures; when this is done, consistent counts can be achieved by pathologists.(van Diest, Baak et al. 1992) After we follow standardized methods, these criticisms should be re-evaluated for manual and automated MC. MC can be determined by partially automated means, using artificial intelligence (AI, more specifically, deep learning-based algorithms). MC performed with computer systems can correct for interobserver variations associated with manual counts. They can better

identify hot spots (Aubreville, Bertram Deep learning algorithms outperform...2020), and 439 they can count thousands of fields but may introduce different hurdles. High quality 440 training datasets that adequately reflect the variability of histopathology sections and 441 scanned images, along with validation of AI methods are paramount for CPATH to 442 produce accurate and verifiable counts. With high quality data sets that define MF, 443 atypical mitotic figures (AMF) possibly along with hard negatives such as mitotic like 444 figures (MLF such as inflammatory cells or cells undergoing necrosis or apoptosis), 445 automated means to perform MC should eventually be able to address potential 446 confounders. Regardless of which mode, manual or automated, we propose that each 447 of these elements needs to be standardized: 1. definition of MF, AMF and MLF; 2. the 448 size of the area in which MF and AMF are counted; 3. the area of the tumor to be 449 evaluated and 4. how to handle confounders. Each of these is described in Appendix 1 450 and 2, CPATH is in Appendix 5. At the end of all appendices are considerations for 451 future studies which should help improve the method and clarify issues associated with 452 assessing the parameter. 453

454 Histologic Morphology of Mitotic Figures (MF) (See Appendix 2)

455 What morphological features define a MF to be included in a MC?

The morphologic characteristics of MF and AMF and features which distinguish these 456 from MLF are detailed in a recent publication. Mitotic figures and AMF are most easily 457 identified by the short "rods" of chromosomes protruding from the surface of aggregates 458 of nuclear material (Figures 1-4). Identification of the different phases of mitosis or the 459 type of AMF are not necessary, but an understanding of the mitotic continuum and that 460 461 AMF may have prognostic significance should be appreciated. Counting AMF may correlate with poorer prognosis and outcome as seen in some human tumors.(Jin, 462 463 Stewenius et al. 2007, Matsuda, Yoshimura et al. 2016) Definitive MF (figures 1-4) and 464 AMF (figures 5-8) should be included in the MC; however, structures with ambiguous morphology create a dilemma in classification. This is not problematic if the MC is 465 markedly high (e.g. >20 MF/2.37mm²). However, if the MC is close to an established 466 threshold which has clinical significance, then the identity of these candidate structures 467 could be critical (see MC Appendix 1.0). New thresholds should be established following 468

the guidelines in Appendices 1 and 2 and those thresholds should be tiered (avoid 469 thresholds based on a< or> single number). Clinicians that request recounts because 470 471 the MC of a tumor is at a threshold should seek different parameters to help establish the prognosis or direct therapy. We all likely have "non-standardized methods" that we 472 use while counting MF but would not necessarily like others to know we do them: 473 enumerating doubtful structures under a column labelled "?"; looking at extra fields 474 when no MF were seen; looking at extra fields because there were spaces created by 475 blood vessels, ducts or cysts; what to do when the tissue sample is <2.37mm²; and/or 476 looking for MF when the diagnosis of inflammation vs neoplasia is not clear. Practical 477 considerations while performing a MC are listed in Appendix 1. Pathologists and 478 laboratories will develop their own procedures to address MC reporting in non-routine 479 situations. When solutions are found, the appendix will be updated accordingly. Correct 480 identification of histologic structures will improve MC consistency and accuracy obtained 481 from manual (glass or WSI) or CPATH modes. 482

483 Does the FN of an ocular matter?

For light microscopy, absolutely. It is the limiting factor that determines the diameter and 484 therefore the area in the field of view (FOV) when objectives of the same magnification 485 are used. Engraved or printed on some ocular eyepieces is a field number (FN) ranging 486 from 6-28 mm. Higher numbers have larger FOV diameters and small increases in the 487 488 FN will produce large increases in the area of FOV (see Appendix 1). The diameter of the FOV can be measured with a stage micrometer or it can be calculated by dividing 489 the FN (mm) by the objective magnification. The formula for the area of a circle is used 490 to calculate the area in the FOV. Therefore, a microscope with an ocular FN 18mm, 40X 491 492 objective has a diameter of 0.45mm in the FOV and an area of 0.16 mm² per "hpf"; FN 26.5mm, 40X objective has a diameter of 0.66 mm and an area of 0.34 mm² per "hpf" 493 which is a 100% larger area, a two fold increase (see Figure 3; Table 3 in Appendix 494 1).(Meuten, Moore et al. 2016) 495

Some objectives will have FN and/or NA (numerical aperture) numbers engraved or
printed on them. Both are defined in the Appendix 9 "definitions and abbreviations". NA
is critical for resolution and depth of field but it is not used to calculate FOV. The higher

the NA the greater the resolution, or sharpness of features. All objectives have an FN
but it may not be engraved on the objective. The FN of an objective can influence the
FOV: however, it is the ocular FN which limits the maximum size of FOV in a standard
microscope, not the objective FN.

503 What is the area in 10 high power fields (hpf)?

504 The area in 10 hpf is not a standard size as it varies up to 200% or more with the 505 objective and the FN of the ocular. (Meuten, Moore et al. 2016) We proposed replacement of the imprecise phrase, 10 hpf with 2.37 mm² to reflect the area equating 506 to 10 hpf using a 40X objective and a 10X ocular FN 22mm, the most common 507 configuration of pathologists' microscopes today. Furthermore, 10 hpf is nebulous for 508 whole slide imaging which is likely the number one means for diagnostic tumor 509 evaluation worldwide. A standard size area in mm² is required so the characteristics of 510 the monitor and the magnification at which the image is reviewed can be configured to a 511 specific area (see Appendix 1). Temporarily, retaining the phrase "10 hpf" together with 512 accurate terminology (2.37 mm²) clarifies communication with clinicians and permits MC 513 to be determined with microscopes or WSI. 514

Mitotic counts (MC) reported in terms of high-power fields (hpf) without specific units of 515 measurement (mm²) cannot be compared to other MC as the area within one or 516 especially 10 hpf is too variable.(Meuten, Moore et al. 2016) Older microscopes were 517 equipped with ocular FN 18 (smaller FOV) compared to current microscopes which 518 commonly have oculars of FN 22 or greater. Most prior animal studies did not define 519 the area (mm²) in which the MC or other histological features were enumerated, or 520 defined the area incorrectly limiting the utility of this data for formulating prognoses for 521 current cases. These studies need to be repeated with standardized methods of 522 523 determining the basic histological parameters used to evaluate tumors. New methods should be considered and all must be correlated with outcome assessments. 524

525 Does the standard area need to remain 2.37mm²?

526 No, it can be changed with data driven results. The total area evaluated can be 527 amended for different tumors or unique situations. e.g. total tissue submitted is

<2.37mm²; cystic tumors etc. Perhaps tumors with low proliferative rates require larger 528 areas to be enumerated (5-10mm²) or perhaps it is the opposite. What might be more 529 530 important than a MC in one spot is what proportion of an entire tumor (or section) has low vs high proliferative rates. Greater than 85% of canine cutaneous MCT are indolent 531 (Kiupel, Webster et al. 2011); perhaps determining the percent of a MCT that is "cold" 532 (few hot spots, or areas of high mitotic activity) will predict how aggressive the tumor is. 533 For canine oral melanoma, it might be the proportion of the tumor that is "hot" which is 534 predictive. We also do not know how many sections of a tumor should be enumerated 535 for the MC to be most predictive? This is true for other histologic parameters as well. 536 These changes require correlating the different methods with known outcomes in many 537 cases to show which method is predictive. Once a method is validated for a tumor type, 538 539 the same size area, same region of the tumor and means to identify MF and AMF need to be validated if we want to compare results between labs or use published cutoffs of 540 541 histologic parameters.

When multiple sections or regions are enumerated, should an average MC be reported 542 543 or the ranges? (Meyer, Cosatto et al. 2009) Various guidelines have been proposed for determining the optimum tumor area for performing the MC in human tumors. Different 544 545 sized areas are recommended to perform MC for different tumors. Some authors recommend counting a series of 5 or more sets of MC and reporting the average. 546 Others report the highest MC. There are a multitude of scenarios that need investigation 547 to change how we determine MC, and CPATH will greatly aid these studies because 548 MC can be performed faster, more consistently, and can be performed over differently 549 sized areas in different regions of the tumors. CPATH can report the proportion of a 550 tumor that is *hot* or *cold*. Manual counts for these types of studies will be laborious. 551 552 Studies using CPATH should also include the standard means of determining the MC and compare the various methodologies to known outcomes. Hopefully, these studies 553 will avoid creating MC cutoffs that are based on a single number (above or below) and 554 develop scoring systems, confidence intervals, and ranges of predictability for MC for 555 different tumors. 556

557 Until data driven results provide new methods, an area equivalent to 2.37mm² should 558 be used for MC and should be reported as mm² rather than stating the FN of the ocular 559 or how the scope is configured.

560 Where in the tumor should the MC be performed?

Presently MF and AMF should be counted in regions of *hot spots* or high mitotic activity 561 562 in viable regions of tumor. It is logical to choose regions of high tumor cell proliferation 563 because the cells in these areas may be more aggressive, they already may have the potential to metastasize or they have a greater opportunity to form a clone with 564 metastatic potential. Until studies report that a different region is more predictive of 565 outcomes, we should adhere to this method. There are no studies in animals that 566 correlate MC determined in different regions with outcomes. Multiple studies in humans 567 and one in dogs have demonstrated variability in the number of MF in different regions 568 of tumors.(Bertram, Aubreville et al. 2020) We know there is heterogeneity of MF 569 distribution in tumors, but we do not know if it matters, and we will not know until there 570 are outcome assessments correlated to methods. Different regions and differently sized 571 areas of different tumor types are used to perform MC in human tumors, and different 572 cutoffs of MC are used to determine prognoses. Similar studies need to be done with 573 animal tumors, and when these are performed, investigators should include newer 574 technologies as well (molecular, CPATH etc., https://www.cap.org/protocols-and-575 576 guidelines/cancer-reporting-tools/cancer-protocol-templates)

577 The periphery of some tumors is the preferred site because this is the invasive front,

578 fixation is better, and there is a higher proliferative rate. A study of human breast

carcinoma reported that the periphery contained more hot spots (using Ki67) than other

regions and percentages of Ki67 positive nuclei obtained at the periphery changed the

prognosis.(Gudlaugsson, Skaland et al. 2012) Other studies in humans reported that

using Ki67 in hot spots, which were not just at the periphery of breast carcinoma,

contributed the most prognostic information as compared to other

methods.(Stålhammar, Robertson et al. 2018) Additionally, a study of canine cutaneous

585 mast cell tumors did not find that the regions of highest mitotic activity were always at

the periphery.(Bertram, Aubreville et al. 2020) Selecting the area of a tumor that is

predictive of outcome(s) or treatments needs to be found for each tumor type. Until 587 those locations are identified. MC should be performed in regions of hot spots. 588 589 However, determination of hot spots by routine light microscopy is subjective and a source of interobserver variation.(van Diest, Baak et al. 1992, Bertram, Aubreville et al. 590 2020) A study with canine MCT and one with canine melanoma showed that 591 pathologists were not as capable of finding the hotspots as compared with computer-592 assisted localization of hot spots.)(Puri, Hoover et al. 2019, Aubreville, Bertram et al. 593 2020) 594

595 *Summary*: Appendix 1 and 2 detail the standard method of performing a MC 596 including: definitions of MF, AMF and MLF, contiguous 2.37 mm² area, hot spot, 597 practical considerations, and future considerations of how the MC can be improved. 598 The present standard means to perform the MC will be modified when data-driven 599 changes necessitate, and the appropriate appendices will subsequently be updated.



600

Figures 1-4: Mitotic Figures (MF) are characterized by dark aggregates of nuclear 601 material with short rods and projections. Figure 1: Prometaphase/metaphase (dense 602 nuclear cluster with short protruding rods). Figure 2: Metaphase with linear equatorial 603 plate of darkly staining nuclear material and short protruding rods and spikes. Inset: 604 Ring form of metaphase with end-on (non-perpendicular) view of the equatorial plate. 605 Figure 3: Anaphase MF with two separate nuclear aggregates with irregular contours 606 607 and short protruding spikes. Figure 4: Telophase MF with aggregates at opposite ends of the cell and formation of a cleavage furrow. 608



- 610
- Figures 5-8: Atypical MF (AMF). Figure 5: Tripolar AMF (more than two spindle poles
- during any stage of mitosis). Figure 6: Asymmetric AMF (unequal sizes of the
- 613 metaphase axes or anaphase poles). Figure 7: AMF with anaphase bridging
- 614 (chromosomes stretching from one pole to the other). Figure 8: Lagging chromosomes
- 615 left behind during anaphase (small dark purple streak in center of cell).
- 616
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665 Lymphovascular Invasion (See Appendix 3)

Neoplastic cell invasion of blood vessels or lymphatics is widely recognized as evidence 666 of tumor aggressiveness and potential malignancy in both humans (Falvo, Catania et al. 667 2005, Mete and Asa 2011) and animals (Goldschmidt, Pena et al. 2011, Rasotto, 668 669 Berlato et al. 2017) but, despite, the importance of this parameter, criteria to definitively 670 identify lymphovascular invasion (LVI) and distinguish from pseudo-vascular invasion or retraction artifact are lacking in the veterinary literature. This lack of stringent, 671 672 standardized criteria may have led to misdiagnosis of LVI in veterinary oncology studies. Assessing LVI with criteria of varying stringency has revealed key insights into 673 674 the biological behavior of human cancers as has the distinction between blood vascular and lymphatic invasion. (Van den Eynden, Van der Auwera et al. 2006, Lin, Zhu et al. 675 676 2010, Mete and Asa 2011) In veterinary medicine, LVI is recognized as a marker of potential tumor malignancy but this parameter has only been extensively evaluated in 677 678 canine and feline mammary tumors(Goldschmidt, Pena et al. 2011, Rasotto, Berlato et al. 2017) without establishment of strict criteria for LVI diagnosis or comparison of blood 679 680 vascular and lymphatic invasion.

Mimickers of LVI, such as pseudo-vascular invasion and retraction artifacts are not 681 adequately addressed in the veterinary literature; images of each can be found in 682 683 Appendix 3 and on the website. Pseudo-vascular invasion is the presence of neoplastic 684 cells within vascular spaces, but the cells are not present because of tumor invasion of 685 vessels. Displacement of neoplastic cells into vessels secondary to manipulation of the neoplasm at the time of biopsy, surgical excision, grossing procedure or tissue 686 sectioning (ie, "floaters") can result in pseudo-vascular invasion.(Van den Eynden, Van 687 688 der Auwera et al. 2006, Mete and Asa 2011) This is also reported for non-neoplastic lesions in the thyroid. (Mete and Asa, 2011) Neoplastic cells may protrude or impinge 689 into adjacent vascular lumens without true invasion in which case endothelial cells cover 690 the surface of the impinging tumor. However, endothelium may also line the surface of 691 neoplastic cells which have invaded through the vascular endothelium but have 692 693 undergone re-endothelialization, necessitating searching for other criteria of LVI to confirm which is the correct interpretation. 694

Distinguishing between these various manifestations of pseudo-vascular invasion and true LVI relies on identification of more robust LVI criteria. The two most definitive criteria used to define LVI in human tumors include: thrombus adherent to intravascular tumor and tumor cells invading through the vessel wall and endothelium. Additional criteria are listed in Appendix 3 along with a complete reference list. (Mete and Asa 2011. These criteria should be used to assess LVI in tumors from animals.

Retraction artifact, another mimicker of LVI, forms an artifactual space surrounding
 tumor foci and can be distinguished from intravascular neoplasia by the absence of an
 endothelial cell lining. Retraction artifact is seen in epithelial tumors in which tumor cells
 retract from surrounding stroma (Figure 5 in Appendix 3).

705 Studies of human breast, thyroid and prostate cancer show widespread metastases are more commonly associated with blood vascular invasion in contrast to lymphatic 706 invasion.(Mete and Asa 2011) Animal tumors may show similar distinctions between 707 blood and lymphatic vascular invasion, warranting detailed descriptions of the type of 708 vessels invaded (ie, if a muscular wall is discerned in the involved vessels) or use of 709 immunohistochemical markers to distinguish blood from lymphatic vessels. A variety of 710 immunohistochemical markers have been used to identify endothelial cells in blood and 711 lymphatic vascular channels in humans and animals (Von Beust, Suter et al. 1988, 712 Sleeckx, Van Brantegem et al. 2013, Wennogle, Priestnall et al. 2019, Fitzgibbons, 713 714 Connolly et al. 2020) Some markers, such as CD31 and Factor VIII related antigen, do not discriminate between lymphatic and blood vascular endothelium, whereas others, 715 such as Lymphatic vessel endothelial receptor 1 (LYVE-1), D2-40 and prospero -716 related homeobox gene-1 (PROX-1) are specific for lymphatic endothelium. (Von Beust, 717 718 Suter et al. 1988, Pusztaszeri, Seelentag et al. 2006, Sleeckx, Van Brantegem et al. 2013, Halsey, Worley et al. 2016, Wennogle, Priestnall et al. 2019, Fitzgibbons, 719 Connolly et al. 2020) Use of IHC endothelial markers has been shown to facilitate 720 identification of LVI in tumors in humans(O'Donnell, Feldman et al. 2008,) and in 721 mammary and plasma cell tumors in dogs. (Sleeckx, Van Brantegem et al. 2013, 722 723 Ehrensing and Craig 2018) Validation of IHC markers and antibodies used to differentiate lymphatic vs blood vessels for the different animal species is a necessity. 724

Although IHC confirms the identity of the vascular structure it does not confirm true LVI and, in fact, is not one of the more stringent criteria of LVI.

Studies of tumor lymphovascular density (LVD) in humans have been correlated with 727 LVI in a number of human tumors.(complete list of references in Appendix 3) LVD is an 728 729 enumeration of lymphatics within a defined area of a tumor and is used as an indicator 730 of lymphangiogenesis and therefore probable lymph node metastasis. Both LVD and LVI are used as predictors of lymph node metastases in human breast cancer, and 731 732 peritumoral lymphatic vessels may be the main route for dissemination of the tumor. Intratumoral microvascular density (IMD), the quantitation of blood vessels 733 734 (number/mm²) in tumors, is used as an indicator of angiogenesis or vasculogenesis and by extension LVI and the ability of a tumor to metastasize. New blood vessels in a tumor 735 736 are required for tumors to grow beyond several millimeters; they are believed to facilitate metastasis and are associated with more aggressive neoplasms in humans 737 738 and animals. Although IMD has been assessed in a number of animal tumors and has been associated with higher grade or more malignant histological features (ie canine: 739 740 soft tissue sarcomas, mammary gland tumors, seminomas, cutaneous squamous cell carcinoma,) and cutaneous mast cell tumors),(full reference listing in Appendix 3) there 741 742 have been no comprehensive studies of intratumoral versus peritumoral vascular density nor associations between IMD and blood vascular or lymphatic vascular 743 invasion in domestic animals. Future veterinary studies comparing intratumoral versus 744 peritumoral microvascular density and correlation with nodal and systemic metastases 745 are warranted. 746

A thorough reassessment of LVI is needed in veterinary oncology with attention to the 747 specific details described in the *appendix LVI* and under future considerations. These 748 studies should use the criteria outlined to determine if LVI is present, especially focusing 749 on the more definitive features: invasion through vessel wall and endothelium and 750 thrombus adherent to the tumor. Studies should include detailed descriptions of criteria 751 used to establish presence of LVI and clarify the importance of lymphatic versus blood 752 753 vascular invasion. Quantitation of blood and lymphatic vessels (IMD, LVD) may benefit from the use of CPATH, and both subjective and quantitative analyses should be 754

- correlated with nodal and systemic metastases and, most importantly, known patient
- outcomes.
- 757
- 758 If individuals have images of true LVI and pseudo-vascular invasion please share them
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- 760
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809

810 Necrosis (See Appendix 4)

811 The extent of tumor necrosis has been correlated with tumor biological behavior and is a parameter used in grading schemes in humans. Tumor necrosis has also been 812 included as a grading scheme parameter in animals, primarily in dogs with STS/STT but 813 814 is also used in other grading schemes (canine primary pulmonary carcinoma). Criteria for determining the percent of tumor necrosis in all species have not been adequately 815 described (Kuntz, Dernell et al. 1997, Coindre 2006) Necrosis within a tumor is often 816 subjectively and vaguely used to suggest a tumor is aggressive. In humans, the percent 817 of tumor necrosis has been determined by estimating the amount seen grossly and 818 histologically, whereas animal studies have not indicated if gross observations were 819 used in combination with histological assessment, or if only histologic assessments 820 were evaluated.(Trojani, Contesso et al. 1984, Kuntz, Dernell et al. 1997, Coindre 2006, 821 McSporran 2009, Vayrynen, Vayrynen et al. 2016, Laurini, Blanke et al. 2017, 822 Dobromylskyj, Richards et al. 2020) Many tumors are larger than a histological section, 823 824 and measuring or estimating percentage of necrosis is more problematic. (Chiang and Oliva 2013) 825

In grading human soft tissue sarcomas, necrosis was found to be one of three 826 827 parameters correlating with patient survival and tumor metastasis, along with tumor differentiation and mitotic count.(Trojani, Contesso et al. 1984) The thresholds for 828 829 scoring necrosis histologically were no necrosis (score 0), less than 50% of tumor 830 necrosis (score 1) and greater than 50% tumor necrosis (score 2) but how a pathologist 831 was to estimate those percentages was not detailed. A grade of two could also be assessed for any neoplasm whose gross appearance was described as "mainly 832 833 necrotic" by a surgeon or pathologist even if no necrosis was seen on the submitted 834 sections.(Trojani, Contesso et al. 1984) We do not recommend this last criterion be adopted for animal tumors, and later authors and grading systems, as well as current 835 College of American Pathologist protocol for assessment of soft tissue tumors in 836 humans, require microscopic confirmation/validation of macroscopic evidence 837 suggesting necrosis. (Coindre 2006, Laurini, Blanke et al. 2017) This brings us to the 838

problems associated with gross interpretation of necrosis (and to a lesser extent, its 839 histologic interpretation). Even for an experienced pathologist, the gross diagnosis of 840 necrosis may be problematic, and most pathologists in veterinary pathology will not see 841 the gross specimen. Areas of edema or exudate may be interpreted as areas of 842 necrosis grossly, and areas of hemorrhage, which are often associated with necrosis, 843 may far exceed the boundaries of actual necrotic tissue. These problems are further 844 compounded by certain histologic lesions such as myxomatous change, cystic space 845 formation, edema, hemorrhage and exudate which can resemble or obscure necrotic 846 areas. Gross/macroscopic assessment of necrosis requires histologic confirmation 847 which, in large tumors, may not be practical for veterinary diagnosticians to submit an 848 adequate number of sections (costs) but should be done in research studies. The 849 850 number of sections examined at trimming and or submitted for histopathology for routine diagnostic cases is likely far fewer in veterinary than human pathology. If gross 851 852 assessment is to be used as a parameter, numerous confounders must be clarified in future studies. This requires documentation of systematic sampling of both necrotic and 853 854 viable tissue during the gross examination and confirmation of necrosis by histological evaluation. Alternatively, we can abandon the use of gross assessment and only use 855 856 light microscopy. This would be straightforward, but if gross assessment of tumor necrosis improves the prognostic utility of grading systems then it would be lost as a 857 858 parameter.

Although it seems obvious that the means to assess various histologic parameters need 859 to be defined prior to implementation, this has not always happened, e.g. the area in 860 which MF were counted was never standardized and the same seems true for percent 861 necrosis. The percent of tumor necrosis in soft tissue mesenchymal tumors/soft tissue 862 863 sarcomas (STT/STS) is included in grading schemes, yet the means to assess necrosis has not been clearly defined or standardized. Was the percent necrosis determined by 864 865 examination of the tumor during gross sectioning, and were areas appearing necrotic confirmed microscopically? Was the percent necrosis used in the grading system based 866 upon visual estimate of necrosis in random histologic tumor sections? Was a consistent 867 portion of the tumor submitted for microscopic examination? A recent publication 868 869 suggested preparation of 1 tissue block for each 2 cm diameter of soft tissue

tumors(Roccabianca, Schulman et al. 2020). Since no formulae for number of
blocks/slides per tumor have been described in published grading systems for dogs this
seems like a good starting point, but no studies using this guidance were referenced.

The necrosis appendix (Appendix 4) provides guidelines for recording and scoring 873 874 extent of tumor necrosis on gross and histologic tumor evaluation which should enable 875 evaluation of the utility of this parameter to assess tumor prognosis and patient outcomes. The scoring system proposed is based on prior reports and is indicated 876 above but includes an unusual percentage of <10% for future studies and explains the 877 logic for this. Additionally for necrosis to be objectively assessed as a parameter for 878 879 future grading schemes, new studies must determine if gross assessment of necrosis can be documented in a standardized fashion and if this parameter correlates with 880 881 outcome assessment independently or as part of a grading system. For this to be accomplished, grossing personnel must include sectioning of tumor sites which appear 882 883 necrotic, hemorrhagic, or edematous, regions typically avoided in most grossing procedures. Most veterinary pathologists will only have microscopic sections to estimate 884 885 necrosis and these sections are likely to be a small percentage of the entire tumor. Furthermore, in many cases, the gross description will be inadequate unless grossing 886 887 personnel are instructed to search and report the percent of the entire tumor that appears necrotic. The usual practice of only sampling viable tissue for histological 888 examination might bias the utility of tumor necrosis as an independent or a component 889 parameter in grading systems. Importantly, the size of the tumor, method of sectioning, 890 891 number of cut surfaces examined grossly and histologically must be documented and at some point, standardized. Based on size of tumor, a recommendation is needed for how 892 many sections should be examined grossly and microscopically. It seems obvious that if 893 894 pathologist A examines 5 histologic sections and pathologist B only 1 section of a tumor with 5cm³ dimensions that the data gathered will not be comparable. 895

896 This brings us to the dilemma of how to currently approach reporting tumor necrosis.

897 Given the lack of established guidance, the pathologist can estimate necrosis either

visually with glass slides, WSI or measure necrosis with annotation software in WSI. If

899 WSI has drawing software, simply outline the entire tumor circumference (X) as well as

the areas of necrosis (Y), followed by calculation of X/Y = % necrosis in one section (Fig
1 Appendix 4). In the absence of software or if using a microscope then visually
estimate with varying magnifications (to confirm areas are indeed necrotic) if the percent
necrosis is <50%>. The range of <50%> seems like a wide target and perhaps that is

sufficient for estimates. We "assume" prior studies that estimated necrosis in canine
 tumors only used histology. But how representative the slide(s) are of overall tumor

- necrosis is unknown and inconsistent sampling of the tumor, purposely avoiding areas
- 907 of necrosis in tissue selection can skew any determination of percent necrosis in
- histologic sections. Given the wide target of greater than or less than 50% necrosis, it
- may be possible to assess this level of necrosis histologically, even with inconsistent
- sampling. However, determining a 10% threshold of necrosis may prove problematic,
- as reported in one study indicating that dogs with tumors with > 10% necrosis were 2.7
- times more likely to die of tumor related causes.(Kuntz, Dernell et al. 1997)
- Future studies can clarify how to determine the percent of tumor necrosis, particularly in
- 914 larger tumors, and establish a standardized means of gross tissue selection for
- histologic examination. Various means of assessing for necrosis in histologic sections
- can be compared and statistically evaluated. Results of standardized assessments for
- tumor necrosis can be compared to outcomes in univariate and multivariate analysis in
- concert with other histologic parameters and prognostic utility determined.
- 919

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946 **Computational pathology (CPATH) (See Appendix 5)**

947 Computational pathology (CPATH) is an umbrella term used to broadly encompass computerized/automated gathering of information on disease in patients.(Abels, 948 Pantanowitz et al. 2019) Although CPATH may use a large variety of information 949 950 sources (raw medical data: histology images, radiology images, gene sequences, 951 clinical records), Appendix 5 focuses only on automated image analysis (AIA) of microscopic tumor images, particularly whole slide images (WSI). When used 952 953 appropriately, CPATH is an exciting tool which uses microscopic images (input data) and automatically produces output information (counts or scores of patterns, 954 955 classification of images etc.). It allows the evaluation of large amounts of tumor data on an unprecedented scale, which is likely to reveal novel trends of prognostic importance. 956 957 As AIA is a relatively new modality of analysis in veterinary pathology with a vast number of relevant methods, this field can be overwhelming with respect to terminology, 958 959 technical aspects, requirements for developing algorithms, performance validation, and implementation strategies. Therefore, the associated appendix aims to give an overview 960 961 of relevant terms, general considerations of CPATH methods and specific recommendations for individual prognostic parameters. Generally, two broad categories 962 963 of AIA approaches are applicable for microscopic tumor prognostication: 1) 964 thresholding-based and 2) advanced data-driven approaches. Thresholding-based algorithms use a set of simple, often programmer-designed image processing steps 965 based on the color information of individual pixels, which are especially useful for 966 scoring immunohistochemical labeling intensity. Data-driven approaches learn to 967 retrieve meaningful patterns from images in order to derive the desired information 968 using artificial intelligence (AI). AI can be used with traditional machine learning 969 970 methods that require "hand-crafted" (by developer) information about relevant features of the pattern, or more sophisticated deep learning methods that autonomously extract 971 relevant features (decision criteria are unknown to developers, "black box"). Deep 972 learning is generally more powerful than traditional machine learning methods, but 973 necessitates larger amounts of data. For histological images, supervised learning (as 974 opposed to unsupervised learning) is a very useful method that learns by "feedback" 975

976 from ground truth labels assigned to the input images. Creating those labels is a very977 time-consuming task and is prone to several biases (see Appendix 5).

Possible applications of AIA for tumor prognostication are seemingly limitless and 978 various benefits of these approaches have been determined in previous 979 980 studies.(Stålhammar, Robertson et al. 2018, Steiner, MacDonald et al. 2018, Aubreville, 981 Bertram et al. 2020) Compared to manual assessment by pathologists, algorithms have higher reproducibility, may have higher accuracy, may increase efficiency of repetitive 982 tasks (such as counting of mitotic figures (MF)), and can carefully assess vast amounts 983 of data per case (every image section of multiple WSIs at high magnification) without 984 985 fatigue. AIA of immunohistochemical labeling intensity was reported to have higher reproducibility and improved prognostic value compared to the manual approach by 986 987 pathologists for Ki-67 index in human breast cancer, (Stålhammar, Robertson et al. 2018) and membrane-binding biomarkers in human esophageal 988 989 adenocarcinomas. (Feuchtinger, Stiehler et al. 2015) An automated topometric segmentation mapping algorithm of immunolabeled MF (anti-phospho-histone H3) was 990 991 used to identify mitotic 'hot spots' in canine melanomas and subsequently used image registration in order to assign the same region to H&E stained tumor sections(Puri, 992 993 Hoover et al. 2019) Deep learning approaches for MF identification in H&E stained 994 tumor sections have been developed for human(Veta, van Diest et al. 2016, Aubreville, Bertram et al. 2020) and canine(Aubreville, Bertram et al. 2020) breast cancer as well 995 as canine mast cell tumors. (Bertram, Aubreville et al. 2019) Deep learning-based 996 997 algorithms are comparable with pathologists for counting MF (in the same tumor regions)(Veta, van Diest et al. 2016) and outperform pathologists in identifying the 'hot 998 spot' regions in WSI.(Aubreville, Bertram et al. 2020) However, correlation of algorithmic 999 1000 MC to patient outcome has not yet been investigated in human and animal tumors. For automated metastasis identification in H&E sections, deep learning-based algorithms 1001 can be used for prescreening of images, and a computer-assisted approach has been 1002 shown to have higher sensitivity and diagnostic speed compared to the unassisted 1003 pathologist. (Steiner, MacDonald et al. 2018) Recent studies on tumors from humans 1004 reported that the systems used could even predict if a tumor was benign, carcinoma in 1005

situ, or invasive carcinoma(Aresta, Araujo et al. 2019) as well as predict genetic
 alterations and gene expression from H&E tumor sections.(Kather, Heij et al. 2020)

Algorithms are not flawless, have multiple sources of error (depending on the 1008 algorithmic approach and available dataset) and therefore require very careful validation 1009 1010 (see Appendix 5). While thresholding-based approaches have high explainability of 1011 algorithmic predictions, data-driven approaches are often considered a "black box" as decision criteria of the algorithms are typically unavailable. Although algorithms are 1012 1013 100% reproducible (same result for the same image using the same model), they may not necessarily cope with variability introduced via biological and pre-analytic factors 1014 1015 (tumor type, tissue types present, section preparation and image acquisition). For example, a deep learning-based algorithm for MF may perform poorly on images 1016 1017 obtained from a WSI scanner that was not used for the training images. (Aubreville, 1018 Bertram et al. 2020) If not part of the training data, algorithms can be compromised by 1019 images with very poor tissue or image quality (artifacts, poor fixation etc.). In contrast to thresholding-based approaches, data-driven algorithms are, however, capable of 1020 1021 learning a certain degree of image variability and training datasets should include realistic variability that reflects the intended use. Performance evaluation should be 1022 1023 done with great care, and data-driven approaches can be assessed by mathematical 1024 evaluation (see Appendix 5), (Abels, Pantanowitz et al. 2019) whereas thresholdingbased approaches are often only assessed visually by a pathologist. (Aeffner, Wilson et 1025 al. 2016) As opposed to pathologists, current algorithms are not capable of modifying 1026 1027 their decision based on surrounding tissue (spatial awareness), which can lead to false detections. For example, pathologists are more careful when classifying a MF in an area 1028 of necrotic tissue as it may be a MLF but algorithms will not use surrounding tissue and 1029 1030 will use the decision criteria programmed to evaluate the candidate structure.

Besides the numerous hurdles in development of AIA algorithms, there are practical
issues to consider for bringing AIA into diagnostic workflows. Basic requirements
include consistent tissue preparation steps, a digital image acquisition workflow,
appropriate IT infrastructure, and sufficient computational power. Increasing
implementation of digital microscopy in veterinary laboratories(Bertram and Klopfleisch)

1036 2017) will augment access to WSI and facilitate AIA. Nevertheless, acceptance of AIA may be hampered by unfamiliarity, limited research results and poor explainability of 1037 1038 machine learning-based algorithms ("black box"). However, there are approaches that can convert the "black box" into a more transparent "glass box" that are likely to have 1039 higher acceptance. For example, some algorithms can be implemented as computer-1040 1041 assisted prognosis systems (as opposed to fully computerized decisions) that always require review by a pathologist. These approaches will improve the reliability of the 1042 computer assisted prognosis system and allow the reviewing pathologist to retain 1043 responsibility in making final decisions with regards to these prognostic parameters. AIA 1044 could greatly improve tumor prognostication by providing vast amounts of reproducible 1045 1046 and possibly accurate information on the tumor section, but interpretation of the result

- 1047 remains the responsibility of the pathologist.
- 1048

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1098 Margins (see Appendix 6)

Margin assessment is one of the most important histological parameters evaluated in oncology.(Kamstock, Ehrhart et al. 2011, Stromberg and Meuten 2017, Liptak 2020) Patient management decisions often hinge on the results of margin assessment, and clinicians may value margin assessment as highly or more than a diagnosis. Appendix 6 provides the types of data that are required to standardize the reporting of margins for both clinical management and future studies.

1105 Histologic margin evaluation only needs to be reported on tumors where the aim of surgery is to completely remove the neoplasm (achieve local control). Samples where 1106 there was no intent to totally excise the tumor, including intralesional or incisional 1107 biopsies are for diagnosis only. Certain tumors or the anatomic location of a tumor 1108 1109 dictate that excision for local control will be attempted but adjacent structures limit how much margin can safely be taken, and therefore margin assessment by the pathologist 1110 should not be requested, e.g. thyroid, anal sac tumors, adrenal glands. Consensus was 1111 not reached whether margins should be reported for benign tumors. Appendix 6 1112 provides contrasting philosophies (*Why not? vs Why bother?*) and the recommendation 1113 that considerations such as this should be left at the discretion of the pathologist and or 1114 1115 their lab as there was no data to support either approach.

For the overall evaluation of surgical margins, the members of the cancer 1116 1117 treatment team are the clinician, surgeon, laboratory technologist and pathologist. The responsibilities of each are detailed in Appendix 6. Although terms such as complete, 1118 clean, clean but close, narrow, and dirty are ingrained in the clinical and pathology 1119 lexicon, practitioners, surgeons, and oncologists should discourage their use and not 1120 1121 expect these to be used in pathology reports. Ultimately it is the clinician and/or surgeon 1122 that judges if the margin is deemed adequate after consideration of all factors. Observations by the pathologist include 1) relationship of neoplastic cells to the 1123 surrounding tissue including presence of a capsule, tissue compression, peripheral 1124 1125 invasion and lymphovascular invasion 2) the distance from neoplastic cells to the narrowest or closest inked margin (histologic tumor-free distance (HTFD, Figure 1) and 1126

3) the relationship of neoplastic cells to the boundaries of the *compartment* in which the 1127 tumor is located. In many cases, measuring the HTFD alone is not enough to determine 1128 1129 the adequacy of surgical margins, yet it is the parameter that is often used to determine 'completeness' of excision by clinician and pathologist. Inking the margin by the 1130 clinician/surgeon immediately after tumor excision is required if a HTFD is expected. 1131 1132 Although surgical margin identification/inking is routinely performed by most surgeons, this practice is not commonplace in general practice. Details of how to apply ink have 1133 been reported (Kamstock, Ehrhart et al. 2011, Appendix 6) and this information should 1134 be included in veterinary school curricula. If ink is not present when the sample arrives 1135 at the lab this should be noted. Only a small portion of the circumferential surgical 1136 margin is evaluated histologically (approximately 0.1-0.01% of the total margin)(Rapini 1137 1138 1990, Becker 2007, Selmic and Ruple 2020). HTFD should be further studied by comparing different methods of margin analysis (radial, tangential, parallel slicing) with 1139 outcome assessments for different tumor types (Milovancev, Townsend et al. 2017, 1140 Dores, Milovancev et al. 2018). Until those studies provide comparative data, radial 1141 1142 sections are recommended. Regardless of the method used, any margin measured histologically may not accurately represent the tumor and its relationship to the normal 1143 1144 surrounding tissue in the patient. It is important to note that HTFD is made on a histopathology specimen that has undergone shrinkage, (ranges reported from 13-50%) 1145 1146 and can underestimate the surgically obtained margins by up to 40%. (Miller and Dark 2014, Upchurch, Klocke et al. 2018) Most of the shrinkage occurs immediately after 1147 1148 removal and prior to fixation.(Clarke, Banks et al. 2014, Miller and Dark 2014, Upchurch, Klocke et al. 2018) The important margin is between neoplastic cells and 1149 1150 "normal tissues" (non-neoplastic) in the patient and this can only be estimated from 1151 histopathology. It is recommended to use whole numbers and ranges when reporting HTFD as reporting distances with decimals implies a level of precision and confidence 1152 that could be misleading. Furthermore, data is accumulating that the biological behavior 1153 of the tumor may be a more important predictor of recurrence than identification of 1154 1155 neoplastic cells at a margin. Certainly, this seems to be the case with low-grade canine MCT and STS/STT. Most low grade MCT do not recur even with tumor cells at the 1156 margin and approximately one-third of high-grade MCTs will recur when the histologic 1157

margins are free of tumor cells.(Donnelly, Mullin et al. 2015) Similarly, for canine soft
tissue tumors/sarcomas, greater than 95% of canine STT do not recur if margins greater
than 1mm are free of neoplastic cells and one study reported that when margins are
less than 1 mm, three of 41 grade 1 tumors (7%), 14 of 41 grade 2 tumors (34%), and 3
out of 4 grade 3 tumors recurred.(McSporran 2009) The biology of the tumor and the
host (immune system, genes) are important factors that influence tumor recurrence and
metastases.

Of importance to surgeons is the concept of compartmental boundaries, which 1165 are used to plan and perform surgical removal of tumors. (Enneking, Spanier et al. 1980, 1166 Kawaguchi, Ahmed et al. 2004) The surgical margins for tumors can be planned 1167 differently if the tumor is in a well-delineated anatomic compartment (such as bone, 1168 joint, muscle) or is infiltrating poorly demarcated interfascial planes and 1169 spaces.(Enneking, Spanier et al. 1980, Kawaguchi, Ahmed et al. 2004) For a well-1170 1171 delineated compartment, it should be reported whether the tumor penetrated the anatomic structure forming the boundary (e.g. periosteum, epimysium or cortical bone). 1172 1173 The deep fascia has been described as a barrier of the subcutaneous tissue compartment(Enneking, Spanier et al. 1980) but this structure is not always included in 1174 1175 sections of cutaneous and subcutaneous tumor resections. Compartment boundaries may be natural barriers to tumor extension. (Enneking, Spanier et al. 1980, Kawaguchi, 1176 1177 Ahmed et al. 2004) It is unclear if these structures function as a true barrier to tumor growth; if they do then it is likely multifactorial and depends upon the aggressiveness of 1178 1179 the neoplasm as well as the components of the barrier (eg cortical bone vs adipose tissue; cytokines). Furthermore, what a surgeon vs a pathologist sees as a *fascial plane* 1180 1181 may not be the same. If pathologists report the facts of what structures were seen 1182 between the tumor and the inked margin, surgeons and oncologists can decide if they are appropriate barriers, and if so, the clinical significance of their presence. Future 1183 studies need to clarify if anatomic structures can prevent tumor infiltration, if so how and 1184 what the pathologist should identify for skin and subcutaneous "tissue barriers" and 1185 1186 fascial planes. (Fulcher, Ludwig et al. 2006) Appendix 6 lists references that describe using CT and MRI for visualizing tissue compartments and assessing the relationship of 1187 1188 tumor to adjacent structures, even differentiating aggressive from benign soft tissue

tumors in humans. It is reported that the tunica serosa fascia in peritoneal cavities is a
barrier to migration of tumor cells using an in vitro system.(Gao, Ye et al. 2013)
When a delineated anatomic compartment is not obvious, the HTFD is of critical

importance. HTFD for lateral and deep margins in samples from skin and subcutis 1192 tumors should be reported separately. In a review of surgical biopsy reports of canine 1193 cutaneous mast cell tumors, details about the margins and consistency of how 1194 histologic margins were reported were generally lacking.(Reagan, Selmic et al. 2018) 1195 For example, while some margins were reported in 92% of cases, lateral and deep 1196 margins were described separately in 77% of cases, margin direction was only given in 1197 16% of cases and descriptions of the deep margin were only available in 11% of 1198 cases.(Reagan, Selmic et al. 2018) The deep margin is difficult for surgeons to visualize 1199 intraoperatively. At the end of appendix 6 are considerations for future studies (M1-M4 1200 or R0-RX)(Stromberg and Meuten 2017, Liptak 2020) 1201



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- 1203

Figure 9: Canine cutaneous mast cell tumor involving the dermis and subcutaneous tissues. The histologic tumor free distance (HTFD) is depicted with horizontal and vertical black lines and can be measured with manual or digital means. Note that ink

- 1207 can be observed at the lateral (or peripheral) margins, but is not visible at the deep
- 1208 margin. Therefore, the deep margin measurement represents an approximation given
- 1209 the lack of ink. Additional sections into the formalin fixed, paraffin embedded block may
- resolve this issue. A potential tissue barrier within the subcutaneous tissue is the
- 1211 striated muscle (also called panniculus carnosus or cutaneous trunci in the truncal
- region, denoted by the asterisks). This muscle is not always visible in histologic sections
- 1213 of cutaneous and subcutaneous tumors; it has variable distribution and continuity in
- 1214 different body regions.(Ahmed, Kulikowska et al. 2019) The subcutaneous fat and loose
- 1215 connective tissue are considered a weak barrier as compared to epimysium,
- 1216 epineurium, or periosteum. The effectiveness of tissue barriers is likely multifactorial
- 1217 and depends upon the aggressiveness of the neoplasm as well as the components of
- 1218 the barrier.

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1277

1278 Outcome Assessment (See Appendix 7)

1279

1280 Standardized methods of histologic and outcome assessment parameters for animal tumors are essential if we wish to compare studies and apply the data to clinical cases. 1281 1282 The protocols and the appendices herein are an attempt to address this goal and 1283 complement existing literature. Outcome assessment of clinical patients is required for determining the predictability of histologically determined prognostic measures (e.g., 1284 tumor classification, grade, etc.) Outcome assessment data need to be collected as 1285 carefully and accurately as the techniques used to assess tumors. (Webster, Dennis et 1286 al. 2011) Some criteria are subjective, clinical, and out of the realm of pathology. 1287 Clinicians must carefully select and standardize clinical outcome measures to avoid 1288 1289 potential confounders. For example, reporting either disease- or progression-free interval is preferable to median survival time, in order to avoid the confounding effect of 1290 timing of euthanasia, which reflects individual biases present within owners and 1291 clinicians. Pathologists play a critical role in accurately determining both progression-1292 and disease-free intervals by allowing definitive determination of whether the same 1293 tumor recurred and/or metastasized given the appropriate tissue. Obtaining samples for 1294 1295 histopathology presents more of a challenge than non-invasive imaging modalities. 1296 Although many advances have been made in this realm, microscopic examination of 1297 tissues remains the gold-standard. Histopathologic assessment has advantages over cytologic evaluation as more definitive information regarding tumor type can be gained 1298 from histopathology. Spindle cell tumors pose a particular problem for cytologic 1299 1300 evaluation as it is difficult (impossible) to distinguish reactive fibroplasia (granulation 1301 tissue) from sarcomas and difficult to specifically identify tumor type. If we want to know if there is reactive fibroplasia or recurrent perivascular wall tumor at the original excision 1302 1303 site, histologic assessment is ideal. However, even with histopathology it is difficult to differentiate these two processes and can be difficult to find tumor cells in re-excision 1304 1305 specimens. There is no standard means to evaluate these cases (clinically and histologically) and in at least one study of STTs, presence or absence of tumor in re-1306

excision specimens did not accurately predict recurrence.(Bacon, Dernell et al. 2007)
Future studies could include imaging modalities, and correlate outcome with the
presence of normal tissue at the margins of resected samples (eg. no spindle cells of
any type). The type of tumor being evaluated will influence the feasibility of visualizing
residual tumor cells in margin excisions as well as the concern for local recurrence.

1312 Genetic studies have shown human and animal breed susceptibilities to develop specific tumor types and multiple tumors in the same host. We know that multiple 1313 1314 aggressive tumors can be present in the same dog, (Golden Retriever, Rottweiler, Bernese Mountain dogs and others)(Cullen and Breen 2016). Given this tendency, it is 1315 1316 essential to make a definitive diagnosis of tumors in metastatic sites. Combining methodologies is ideal but practical considerations of costs and emotional factors 1317 1318 impact study results. Imaging can provide an alternative means to assess for suspected 1319 metastatic lesions and can provide useful clinical information for patient management 1320 but leaves a gap in outcome assessment studies provided no other confirmatory data is available. Imaging cannot determine whether the tumor suspected of being in the lungs 1321 1322 is the same tumor as was excised previously. These methods to identify suspected neoplasia are the methods of choice for clinical settings but not research models. It is 1323 1324 important to differentiate the information from a test being used to help treat one patient 1325 or predict how a population of animals with the same tumor will react to that tumor type. The latter will be applied to the former when we gather and analyze data carefully. 1326 Histopathology remains the gold standard to develop ground truths if the tumor type is 1327 the same. We can substitute other methods for histopathology, but the data should be 1328 labelled suspected neoplasia/metastases (e.g., as determined by imaging or physical 1329 exam) but not confirmed unless histopathology is used. In the future, molecular testing 1330 1331 of suspected tumor tissue may be superior to histopathology.

Other appendices have detailed how to assess parameters used to evaluate a tumor, recurrence, margins and metastases. In order to use morphologic diagnoses, margins, LVI, MC, lymph node status, or CPATH to predict tumor behavior and/or to select treatment options, oncologists must acquire sufficient outcome assessment information to allow interpretation of tumor parameters. Knowing actual survival times of geriatric

pets or including pets in which no treatments were performed provides control groups to 1337 which treatments and outcomes can be compared. Determining the least invasive 1338 1339 means to characterize tumor behavior is ideal but cannot be accomplished without adequate outcome assessment studies. Appendix 7 utilizes and expands upon 1340 published guidelines for conduct and evaluation of prognostic studies(Webster, Dennis 1341 1342 et al. 2011) and for response assessment in canine solid tumors, (Nguyen, Thamm et al. 2015) citing specific information gained from studies of canine soft tissue sarcoma and 1343 canine mast cell tumor. 1344

Standardized criteria, such as RECIST and RECIST 1.1(Therasse, Arbuck et al. 2000, 1345 1346 Schwartz, Seymour et al. 2016) should be used to document the patient's response to treatment and progressive disease. The RECIST 1.1 criteria have been robustly 1347 1348 evaluated for use in human clinical trials and can be easily adapted to the evaluation of 1349 veterinary patients. Pathologists, oncologists, surgeons, clinicians and students should 1350 be familiar with the terms explained in these manuscripts which indicate response to treatment and include Complete remission (CR), Partial response (PR), Progressive 1351 1352 disease (PD), Stable disease (SD) and Not evaluable (NE).(Nguyen, Thamm et al. 2015) Documented progression is needed in the cases of questionable lesions, or a 1353 1354 minimum size is required to determine whether neoplastic disease is present within a 1355 lymph node. Additionally, there may be specific anatomical locations evaluated depending on the tumor type. For example, prostate cancer may favor bone 1356 metastases, pulmonary carcinoma in cats requires assessment of all digits, and 1357 hemangiosarcoma is the most common metastatic tumor to the brain of dogs. Ideally, 1358 imaging will be used in concert with biopsy or autopsy in order to confirm recurrence 1359 1360 and metastasis with the utmost accuracy.

Metastasis should be subdivided into confirmed and suspected. Metastases determined by imaging only should be labelled suspected. Histopathology is required to confirm metastases are present and are of the same tumor type. The preferred methodology of evaluation in humans, the CT scan, should be used if possible as it avoids some of the technical problems associated with the use of radiographs, whereas ultrasound is not an acceptable method of assessing disease state(Nguyen, Thamm et al. 2015) The use 1367 of functional imaging (PET scans) is increasingly common to better determine sites of

- disease; however, it cannot be used for measuring purposes. Following these
- 1369 standardized criteria will ensure that studies can be reproduced and compared between
- institutions, resulting in more useful correlates of clinical data to prognostic information,
- and ensuring progress in veterinary oncologic pathology.
- 1372 Euthanasia is a reality of veterinary medicine, and oncology studies that use pets must
- 1373 carefully evaluate how decisions to euthanize influenced survival times. Reported
- 1374 patient survival times are impacted by euthanasia which may be elected due to
- 1375 perceived pet value, owner income, primary vs referral centers or other factors which do
- not reflect tumor behavior. When patients are euthanized, clinicians should determine
- 1377 and/or record the cause of death with as much accuracy as possible. If euthanasia is
- due to an unrelated disease process, this must be noted. If euthanasia is caused by the
- 1379 neoplasm being studied, and cachexia is present, then histologic confirmation of the
- extent of the neoplastic disease helps verify clinical observations and reliability of study
- 1381 conclusions. Oncology studies no longer include results of autopsy, the perceived value
- 1382 of which seems to have hit a nadir. Permission to perform autopsies should be pursued
- 1383 as autopsy greatly increases the confidence in results from the case. Studies should set
- a goal of autopsies on at least 20% of the cases.
- 1385

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1410

1411 Synoptic Reporting in Veterinary Medicine (See Appendix 8)

1412

Synoptic reporting (as opposed to the traditional narrative reporting) is a method for 1413 1414 reporting specific pieces of prognostically-relevant data in a discrete format in pathology 1415 reports (Renshaw, Mena-Allauca et al. 2018). In human medicine, these have progressed from individual efforts (Markel and Hirsch 1991) to being mandated by the 1416 1417 College of American Pathologists (CAP) for accreditation (College of American Pathologists 2020). In general, a synoptic pathology report consists of data elements 1418 1419 and responses (see Figures S1, S2 Appendix 8, supplemental), which may be either required or optional. For CAP purposes, the report must have all required core 1420 1421 components reported, all conditional core components reported when applicable, must be listed with the element next to its associated response, and all elements and 1422 1423 responses must be on separate lines and in one place in a report. Synoptic reporting has been shown to make pathology reports more readable to clinicians and patients 1424 1425 (Renshaw, Mena-Allauca et al. 2018), as well as making reports more likely to include all data elements needed (Karim, van den Berg et al. 2008, Kang, Devine et al. 2009, 1426 1427 Srigley, McGowan et al. 2009, Messenger, McLeod et al. 2011) To develop an effective synoptic report typically requires the efforts of pathologists and clinicians, who develop 1428 the checklist of required and recommended items after reviewing the relevant literature 1429 (Chamberlain, Wenckebach et al. 2000). Currently, there are two main groups 1430 producing templates in human medicine, CAP and the International Collaboration on 1431 Cancer Reporting (ICCR). Both require a committee of pathologists, oncologists, and 1432 other interested representatives (e.g., World Health Organization working groups, etc.) 1433 1434 to develop a new protocol.

A number of studies have found that synoptic reporting produces reports that are more likely to contain all significant pieces of information than narrative reports. For pancreatic tumors, 100% of synoptic reports had information about small vessel and perineural invasion, compared to 66% and 84% of narrative reports, respectively (Gill, Johns et al. 2009). In addition, the stage could be determined in 100% of synoptic reports compared to 56% of narrative reports. In a comparison of melanoma reports,
mitotic count, histologic subtype, predominant cell type, vascular and lymphatic
invasion, neurotropism, desmoplasia, and distance to the nearest margin were all
reported significantly more frequently in synoptic reports than narrative reports, both at
the teaching institution responsible for the study and the outside reports sent in to the
teaching institution for a second opinion(Karim, van den Berg et al. 2008).

While full implementation of standardized reporting would allow for easy automated data 1446 1447 collection(Ellis and Srigley 2016), even simple implementations of synoptic reporting can allow for significant automated information extraction. For example, if all deep 1448 1449 margins are listed as "DEEP MARGIN: <xx>mm" on a line by itself, it is comparatively easy to extract all margins from reports using standard text search and manipulation 1450 1451 tools (e.g., grep, cut, etc.). Not only can this improve retrospective studies, but can also 1452 provide valuable clinical information, as extracted information can be compared 1453 between services, clinicians, and other variables to determine if these influence patient 1454 outcomes.

1455 From the beginning of synoptic reporting, clinicians have reported increased satisfaction with synoptic vs. narrative reports (Markel and Hirsch 1991). A study of treating 1456 physicians and pathologists in Canada found that both groups found synoptic reports 1457 1458 easier to find information in, facilitate a consistent approach to interpretation of 1459 diagnostic and prognostic factors, and provide higher overall satisfaction (Lankshear, 1460 Srigley et al. 2013). While pathologists felt that reports took approximately 25-50% longer to complete, treating physicians did not notice a difference in the length of time it 1461 took pathology reports to be completed. 1462

The major problem in veterinary medicine is a lack of knowledge about factors involved in prognosis. As discussed in the other appendices in this document, there is little standardization of methods used in determining prognostic factors. There are also no standards for terminology, such as immunohistochemical findings (e.g., "positive" vs. "immunoreactive" vs. "present"), which hinders design of standardized reports. Another issue for many pathologists, particularly in academia, is the effect switching to synoptic reports would have on resident training. Given the necessity of writing descriptions for boards and the lack of universal adoption of synoptic reporting, residents still require
significant experience in writing narrative reports. This can be mitigated by requiring
narrative reports in other resident educational settings (such as rounds) to provide
practice in writing narrative reports for neoplasms.

1474 Many pathologists are concerned about increased time to finish reports with synoptic 1475 reporting, including physicians (Lankshear, Srigley et al. 2013); however, when synoptic reports have been implemented many of these concerns have been deemed 1476 1477 technology related rather than issues with the reporting format. As with many new processes, we assume that once the pathologists become familiarized with the new 1478 1479 format, there will be a decrease in time to write these types of reports. A standardized formatted template will be created and added to the website we propose. In veterinary 1480 1481 medicine, no current laboratory information management system (LIMS) can use synoptic reporting, which may seem like an obstacle to implementation of synoptic 1482 1483 reporting. However, any word processor can be used to implement synoptic reporting without specialized software(Ellis and Srigley 2016); all that is required is to type the 1484 1485 data element, a separator (such as TAB), and the response. Templates can be saved 1486 containing required and optional data elements, making it easier for pathologists to fill 1487 out reports quickly. These can then be copied and pasted into any LIMS or word 1488 processor for subsequent reporting.

Finally, another major obstacle to implementation of synoptic reporting is a lack of awareness of synoptic reporting and its benefits in veterinary medicine. Establishing working groups with pathologists and oncologists to develop guidelines for specific neoplasms would help promote awareness and develop reporting checklists that would benefit both pathologists and treating clinicians.

The next step beyond synoptic reporting is standardized reporting, that is, having a standardized, specific set of responses for each required question(Srigley, McGowan et al. 2009). Ultimately, this can lead to automated staging and grading, as well as improving data harvesting for future research and clinical applications. The addition of free text fields associated with standardized options would allow for customization of reports while retaining standardization for further applications.

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- 1530

	Synoptic Report	
	MASS SIZE:	3 cm x 2 cm x 2 cm
		Gross measurement
	BIOPSY TYPE:	Excisional
	LOCATION:	Forelimb proximal to elbow
	ASSESSMENT METHOD:	Manual light microscopy with glass slides
	HISTOLOGIC TYPE:	Nerve sheath tumor
	DEEPEST LAYER INFILTRATED:	Dermis
		Via histology
	DIFFERENTIATION SCORE:	2
	MITOTIC COUNT:	12 per 2.37mm ²
	NECROSIS:	11-50%
	TOTAL SCORE:	5
	HISTOLOGIC GRADE:	2
		Kuntz system
	LYMPHOVASCULAR INVASION:	None
	METASTASIS:	None
	MARGINS INKED:	By laboratory
	MARGIN TYPE:	Radial
	DEEP MARGIN:	Complete
	DEEP MARGIN HTFD:	3 mm
	LATERAL MARGIN:	Complete
	LATERAL MARGIN HTFD:	6 mm
Į		

Narrative Report

In one transverse and two longitudinal sections (from a 3 x 2 x 2cm mass from the left forelimb, per submitter), the dermis is disrupted by a highly cellular, infiltrative, unencapsulated mass. The mass is composed of cells forming bundles and whorls surrounding empty capillaries. The cells have indistinct borders and eosinophilic cytoplasm. The nuclei are medium to large and fusiform, with finely stippled chromatin. Mitoses average 12 per ten 400x fields (2.37mm²). The central 30% of the mass is necrotic. The mass is separated from the deep and lateral sample margins by 3mm and 6mm, respectively.

DIAGNOSIS:

Nerve sheath tumor, grade II, left forelimb

1531

- 1532 Figure 10: Comparison of synoptic and narrative reports. The same information in each
- 1533 report is in the same color.

1535 Skin and Subcutaneous Soft Tissue Tumors (STT/STS) (See Protocol 1)

1536 This protocol is intended for use with soft tissue tumors arising in the skin and subcutaneous tissues which are predominantly of mesenchymal tissue origin and which 1537 are commonly referred to as soft tissue sarcomas (STS). (Bostock and Dye 1980, Kuntz, 1538 1539 Dernell et al. 1997, McSporran 2009, Roccabianca, Schulman et al. 2020) Modifying a 1540 name generally meets with resistance and lack of unanimity. The term sarcoma suggests the group of neoplasms are aggressive (malignant), however present outcome 1541 1542 assessment data does not indicate that is the case. (Bostock and Dye 1980, Kuntz, Dernell et al. 1997, McSporran 2009, Roccabianca, Schulman et al. 2020) Thus, it is 1543 1544 proposed to remove sarcoma from the acronym. These neoplasms are predominantly mesenchymal, however, a subset (namely nerve sheath tumors) are not solely derived 1545 1546 from the mesoderm, therefore, soft tissue mesenchymal tumor is not entirely accurate. 1547 These neoplasms can be accurately encompassed by the term soft tissue tumors (STT) 1548 (which is admittedly vague), however, ensures that more users of this term will be satisfied. The purpose of this protocol is to provide standards for accruing data so that, 1549 1550 over time, large data sets with comparable information can be evaluated to enable meaningful conclusions and accurate prognostic information. 1551

The term STT/STS encompasses a wide range of benign and malignant tumor types in 1552 1553 humans (Byerly S, Chopra S, Nassif NA et al, 2016) The different types are much more limited in animals and, although the veterinary terminology and various grading 1554 1555 schemes have, in many instances, been borrowed from the human literature, the types of neoplasms which commonly comprise soft tissue tumors in humans are very different 1556 from the tumor types typically encountered in animals. This is exemplified by 1557 1558 liposarcomas, which are common in humans and rare in dogs, and perivascular wall 1559 tumors (PWT), very common in dogs, are rare in humans. Furthermore, STS in humans have extensive molecular profiles to help subtype them, which is not established for 1560 1561 canine tumors. The common denominators between species appears to be an origin in non-epithelial, extraskeletal soft tissues exclusive of hematopoietic system. (Bostock 1562 1563 and Dye 1980, Trojani, Contesso et al. 1984, Kuntz, Dernell et al. 1997, Coindre 2006, McSporran 2009, Dennis, McSporran et al. 2011, Roccabianca, Schulman et al. 2020) 1564

1565 This protocol is intended for use with the following types of tumors: Perivascular wall

tumors (PWT), nerve sheath tumors (NST), fibrosarcoma, myxosarcoma,

1567 leiomyosarcoma, liposarcoma, rhabdomyosarcoma or unclassified spindle cell

tumor/sarcoma arising in the dermis or subcutis. PWT and NST are the most common

1569 types of STT/STS and their biological behavior is primarily indolent.(Roccabianca,

1570 Schulman et al. 2020)The effect of grouping of disparate tumors within the same

1571 grading scheme needs to be compared to grading tumors segmented into specific

1572 histological diagnoses so that important predictive parameters may be determined.

The current scheme used for grading dog STT/STS is patterned after Trojani's grading 1573 1574 of human STS. (Trojani, Contesso et al. 1984) Unlike the human grading scheme, however, the studies of dog STT/STS only evaluated three histological features. Some 1575 1576 criteria, such as determination of the percentage of necrosis via gross and/or 1577 histological criteria, are poorly defined in the human literature and were not clarified in 1578 the veterinary manuscripts. (Bostock and Dye 1980, Kuntz, Dernell et al. 1997, McSporran 2009) Percent necrosis for human tumors was determined by estimating the 1579 1580 amount seen grossly and histologically (see Appendix 4). (Trojani, Contesso et al. 1984, Coindre 2006, Rubin, Cooper et al. 2010, Nguyen, Thamm et al. 2015) There are a 1581 1582 number of distinctions between the grading systems used for human tumors and how 1583 the they are applied to dogs, which have not been addressed in the canine papers; (Bostock and Dye 1980, Trojani, Contesso et al. 1984, Kuntz, Dernell et al. 1584 1997, Coindre 2006, McSporran 2009) in particular, the need to determine histological 1585 tumor type and confirmation of the diagnosis of sarcoma *prior* to applying the human 1586 grading systems. Four additional histological features evaluated by Trojani but not 1587 found useful for human tumors were not assessed in the dog STT/STS grading studies. 1588 1589 Our existing scheme needs to be broadened to determine if parameters originally rejected for human STS may, in fact, be predictive in dogs. The methods described to 1590 assign scores for necrosis, MC and differentiation for canine tumors are not detailed 1591 enough that others can replicate them, and the number of dogs reported with high-1592 grade STT/STS that have outcome assessments is small. These studies need to be 1593 repeated with additional parameters evaluated, more detailed description of methods 1594 1595 and greater case numbers paired with standardized outcome assessments. The

protocol in this appendix provides details of the histological findings that should be
noted in STT/STS which will enable more thorough assessment of these tumors and
should provide a database for performance of studies and validation of grading
schemes.

1600 For any proposed veterinary tumor grading system, the tumor type should be 1601 designated as precisely as possible and the criteria used to designate that diagnosis be provided (H&E, IHC etc). Each graded element must be clearly defined. For instance, 1602 1603 the means to assess percent necrosis (gross, histology, both; Appendix 4) must be clarified if this is an element of a grading system and others are expected to duplicate 1604 1605 the method.(Kuntz, Dernell et al. 1997) Histologic classification of some types of STT/STS is difficult. A particular conundrum is differentiating PWT from NST. 1606 1607 Histological features characteristic of PWT and NST have been described, but there is 1608 overlap of histological patterns found in these two tumor types(Avallone, Helmbold et al. 1609 2007, Suzuki, Uchida et al. 2014, Loures, Conceição et al. 2019, Vučićević, Marinković et al. 2019, Avallone, Stefanello et al. 2020, Roccabianca, Schulman et al. 2020) which 1610 1611 can complicate definitive diagnosis in routinely stained sections. How specific can, or should our diagnoses be from HE slides and how does this influence differentiation 1612 1613 scores used to grade these tumors? Examples: Should PWT be subtyped, and similarly as NST is not just one tumor, should neurofibroma, Schwannoma and malignant NST 1614 be identified? Classification of some tumors, including some cases of PWT, may 1615 require IHC or other ancillary tests. In veterinary medicine, the costs for these tests are 1616 incurred by owners and, if the tests are declined, it is unreasonable to expect a precise 1617 classification of some of these tumors with H&E. These practical factors influence our 1618 diagnoses and grading systems. 1619

Present canine studies have not determined if identifying tumor type is predictive of tumor behavior. Until we use a grading system for specific tumor types as well as for the entire group of STT/STS, we will not know which approach is more predictive. A grading scheme that can be applied to any tumor within the STT/STS group is easier to apply then requiring identification of the specific tumor type before grading, particularly in instances in which a definitive diagnosis cannot be made with evaluation of routinely

stained tissue sections. However, future studies should validate if this is "best practice". 1626 On the one hand, identifying the precise tumor type may have prognostic information 1627 1628 unrelated to a specific grade and, on the other hand, perhaps all tumors within either the entire STT/STS group or within specified subsets of the group (for instance PWT/NST) 1629 may behave according to assigned grades. For instance, group PWT and NST 1630 1631 together, based on H&E histologic morphology and determine outcome and determine if there is a difference in outcome assessment if these two tumor types are evaluated 1632 1633 separately. Identification of these two tumor types may involve IHC or electron microsocopy. If the biological behavior of these two tumors was such that distinguishing 1634 them at the H&E level was not needed that would have practical use for a diagnostic 1635 pathologist and oncologist. The only means to determine the prognostic utility of 1636 1637 grouping or separating tumor types within the STT/STS category is to perform studies which evaluate outcomes related to the STT/STS group as a whole and ALSO evaluate 1638 outcomes in relation to specific histologic type of tumors. Studies must have sufficient 1639 numbers of animals within each tumor grade to generate statistically significant findings. 1640 1641 This latter issue will be a problem for uncommon tumors, such as liposarcoma, for which it may be problematic to find enough high-grade tumors with accurate outcome 1642 1643 assessments, but using criteria in which two tumor types (ie, PWT and NST) comprise more than 80% of the cases to predict how uncommon tumors behave needs to be 1644 1645 validated.

Future considerations should address existing and new grading systems for STT/STS 1646 (see protocol 1). The present grading system should be followed with methods 1647 described in sufficient detail to permit other investigators to duplicate the methods and 1648 the scoring systems. Consideration should be given to assessment of weighted scores 1649 for parameters, such as differentiation or mitotic count, in determining grade and 1650 correlation with outcome assessment. Additional histological features should be 1651 1652 evaluated for their prognostic utility, for instance, tumor cellularity, presence of atypical 1653 nuclei or multinucleated giant cells and presence of lymphovascular invasion (see Appendix 3). The benefit of applying a new, better-detailed scoring system for 1654 1655 histological differentiation should be assessed as this is the most subjective parameter

- in human tumors and likely canine tumors. The use of a defined area in mm² should be 1656
- applied to all parameters enumerated in a grading system. New grading systems 1657
- 1658 should be compared to older systems, and there must be sufficient numbers of animals
- in each tumor grade to enable interpretation of results. Studies should be initiated to 1659
- assess the criteria for diagnosis of NST and PWT and the reproducibility of the criteria. 1660
- Finally, the use of computational pathology and molecular profiling should be explored 1661
- 1662 in determining grades and outcomes of STT/STS.
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