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(Article begins on next page)

Design-based stereological study of the guinea-pig (Cavia

2	porcellus) cerebellum
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4	Running title: Guinea pig cerebellum stereology
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Abstract

22	Guinea pigs have proved useful as experimental animal models in
23	studying cerebellar anatomical and structural alterations in human
24	neurological disease; however, they are also currently acquiring increasing
25	veterinary interest as companion animals. The morphometric features of the
26	normal cerebellum in guinea pigs have not been previously investigated
27	using stereology. The objective of the present work was to establish normal
28	volumetric and quantitative stereological parameters for cerebellar tissues
29	in guinea pigs, by means of unbiased design-based stereology. Cerebellar
30	total volume, grey and white matter volume fractions, molecular and
31	granular layers volume fractions, cerebellar surface area, Purkinje cellular
32	and nuclear volumes, and the Purkinje cell total count were stereologically
33	estimated. For this purpose, cerebellar hemispheres from six adult male
34	guinea pigs were employed. Isotropic, uniform random sections were
35	obtained by applying the orientator method, and subsequently processed for
36	light microscopy. The cerebellar total volume, the white and grey matter
37	volume fractions, and the molecular and granular layer volumes were
38	estimated using the Cavalieri's principle and the point counting system.
39	The cerebellar surface area was estimated through the use of test lines;
40	Purkinje cellular and nuclear volumes were analysed using the nucleator
41	technique, whereas the Purkinje cell total count was obtained by means of

the optical disector technique. The mean \pm standard deviation (SD) total

volume of a guinea-pig cerebellar hemisphere was 0.11±0.01 cm³. The

44 mean volumetric proportions occupied by the grey and white matters were,

respectively, 78.0±2.6% and 22.0±2.6%, whereas their mean absolute

volumes were found to be 0.21 ± 0.02 cm³ and 0.059 ± 0.006 cm³. The

volumes of the molecular and granular layers were estimated at 112.4±20.6

mm³ and 104.4±7.3 mm³, whereas their mean thicknesses were calculated

to be 0.184 ± 0.020 mm and 0.17 ± 0.02 mm. The molecular and granular

layers accounted for 40.7 ± 3.9 % and 37.4 ± 1.8 % of total cerebellar

volume, respectively. The surface area of the cerebellum measured 611.4 \pm

52 96.8 mm². Purkinje cells with a cellular volume of 3210.1 μ m³ and with a

nuclear volume of 470.9 μm³ had a higher incidence of occurrence. The

mean total number of Purkinje cells for a cerebellar hemisphere was

calculated to be $253,090 \pm 34,754$. The morphometric data emerging from

the present study provide a set of reference data which might prove

valuable as basic anatomical contribution for practical applications in

veterinary neurology.

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Keywords: Guinea pig, cerebellum, stereology, neuroanatomy, nervous

61 system.

Introduction

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The involvement of the cerebellum in motor coordination, balance 64 65 and motor learning has been long and widely recognized (Brooks, 1984; Llinás and Welsh, 1993; Baillieux et al., 2008; Lee et al., 2015); however, a 66 growing body of evidence involving neuroanatomical, neuroimaging and 67 clinical studies indicates that it plays a significant role in non-motor 68 behavioral-affective and cognitive functions, as well (Schmahmann and 69 Caplan, 2006; Booth et al., 2007; Molinari et al., 2008; Cantalupo and 70 Hopkins, 2010; Koziol et al., 2011; De Smet et al., 2013; Roostaei et al., 71 72 2014). Design-based stereological techniques allow to efficiently acquire 73 precise quantitative estimates of three-dimensional 74 accurate and morphometric features of whole organs from measurements made on two-75 dimensional sections, by making use of statistical sampling and stochastic 76 77 geometry principles (Boyce et al., 2010). Most stereological investigations on the cerebellum involving 78 laboratory animals have been carried out on mice (Woodruff-Pak, 2006; 79 Woodruff-Pak et al., 2010; Wittmann and McLennan, 2011; Kennard et al., 80 2013; Song et al., 2014), rats (Korbo et al., 1993; Larsen et al., 1993, 2000; 81 82 Ragbetli et al., 2007; Sonmez et al., 2010) and rabbits (Akosman et al., 2011; Selçuk and Tıpırdamaz, 2020), but also on domestic animals such as 83

cats (Sadeghinezhad *et al.*, 2020), pigs (Jelsing *et al.*, 2006) and chicks (Tunç *et al.*, 2006). Apart from a stereological study performed on prenatal and neonatal guinea-pig cerebella following experimentally-induced intrauterine growth restriction (Mallard *et al.*, 2000), the morphometric features of the normal cerebellum in adult animals of this species have not been previously investigated using stereological techniques.

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Guinea pigs (Cavia porcellus) have proved useful as experimental 90 animal models in studying cerebellar anatomical and structural alterations 91 in human neurological disease (Lev-Ram et al., 1993; Furuoka et al., 2011; 92 93 Čapo et al., 2015; Bennet et al., 2017; Cumberland et al., 2017), partly due to their high degree of neurological maturity at birth in relation to the short 94 gestation period (Altman and Das, 1967; Hargaden and Singer, 2012; Silva 95 et al., 2016), which is important for clinical studies in human medicine. 96 Indeed, the brain of newborn guinea pigs is singularly mature, and 97 98 postnatal cerebellar neurogenesis is minimal in this precocial species 99 (Altman and Das, 1967). It was observed that, as early as 45 days through 100 gestation, cerebellar layers in guinea-pig fetuses were distinct and well developed, with easily identifiable Purkinje cells, and with the white and 101 gray matters well differentiated both macro- and microscopically (Silva et 102 103 al., 2016). Moreover, cellular proliferation events in the cerebellum, unlike other rodents, are complete at birth in the guinea pig (Lossi et al., 1997). 104

Recently, however, increasing interest has been addressed toward the clinical features, pathological changes and therapeutic resolution of neurological disorders of guinea pigs held as pet animals (Hollamby, 2009; Hawkins and Bishop, 2012). Most incidences of naturally-occurring cerebellar pathology reported in the literature for pet guinea pigs have an infectious etiology. Reported aetiological agents are, for instance, the lymphocitic choriomeningitis virus, leading to cerebellar hypoplasia with acute destruction of cortex folia and necrosis of granule and Purkinje cells (Monjan et al., 1971; Hawkins and Bishop, 2012); Toxoplasma gondii, inducing granulomatous meningoencephalitis, foci of necrosis, and chronic cysts in the central nervous system (Brabb et al., 2012; Gentz and Carpenter, 2012); and *Bayilisascaris procionis* larvae, causing progressive multifocal encephalomalacia and eosinophilic granulomatous inflammation of the cerebellum, midbrain and brainstem (Van Andel et al., 1995).

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In light of the above-listed scientific evidence, the objective of the present work was to establish normal volumetric and quantitative stereological parameters for cerebellar tissues in adult guinea pigs, by means of unbiased design-based stereology (Gundersen and Jensen, 1987; West, 1993; Boyce *et al.*, 2010). Specifically, the present study was designed to estimate cerebellar total volume, grey and white matter volume fractions, molecular and granular layers volume fractions [by using the

Cavalieri 's principle (Gundersen and Jensen, 1987)], estimate the cerebellar surface area (Schmitz and Hof, 2005), the total number of Purkinje cells [by employing the optical disector method (Gundersen, 1977; Sterio, 1984)], and the mean Purkinje cellular and nuclear volumes [through the use of the nucleator method (Gundersen *et al.*, 1988b)] in the guinea pig.

The morphometric data emerging from the present study provide an accurate set of reference data potentially valuable as basic anatomical contribution to the field of veterinary neurology in order to help implementing the development of the diagnosis and treatment of nervous diseases in the guinea pig.

Methods

Animals and tissue preparation

Six adult male pet guinea pigs, weighing 569 ± 64.9 g, which spontaneously died of diseases other than those affecting the nervous system, were used for our research purposes following owners' permission. The animals did not present a history of neurological disease nor displayed pathological alterations of nervous tissues.

According to Directive 2010/63/EU of the European Parliament and of the 22 September 2010 Council on the protection of animals used for

scientific purposes, the Italian legislation (D. Lgs. n. 26/2014) does not require any approval by competent authorities or ethical committees, as this research did not influence any therapeutic decisions.

Guinea-pig cerebella were excised from the neurocranium in their entirety, each was divided into two halves, and then immersed in a 4% phosphate-buffered formaldehyde solution to enable tissue fixation. One hemisphere of each cerebellum was randomly chosen and weighed on a digital laboratory scale. The cerebellar hemispheres were routinely processed for light microscopic examination and subsequently embedded in paraffin.

Tissue sampling and stereology

The orientator method (Mattfeldt *et al.*, 1990; Nyengaard, 1999) was applied to obtain isotropic, uniform, random sections. In essence, each cerebellar hemisphere was embedded in a paraffin block, which was placed at the center of a circle with 90 equidistant divisions along the perimeter. A random number between 0 and 90 was looked up and the paraffin medium was cut along a line parallel to the direction of the selected number. The block was placed on its cut surface at the center of a second circle, with 96 nonequidistant divisions along its perimeter. The paraffin was cut along a line parallel to the direction of a random number ranging from 0 to 96, and the block was finally re-embedded in paraffin while placed on its cut

surface (Fig. 1). Consecutive 25 micrometer-thick sections were cut with a microtome at uniform constant intervals with a random start and until exhausting the organ. Every 25th section was collected using the principle of systematic uniform random sampling (Gundersen and Jensen, 1987), in order to acquire 12 to 15 sections per animal. Sections were then stained with Cresyl violet 0.1 % stain solution. A slide scanner (Optic lab H850, Plustek) was employed for capturing images from sections in order to enable the subsequent estimation of volumes and surface areas. A microscope (CX40, Olympus, Germany) equipped with an oil immersion objective (×100), connected to a microcator (MT12, Heidenhain, Traunreut, Germany) and a digital camera (MB-225) was utilized for the estimation of Purkinje cells total cellular and nuclear volumes. Geometrical probes, necessary for the stereological analysis of each structural feature represented in each section (West, 1993), were produced using a dedicated software (ImageJ; https://imagej.nih.gov).

Estimation of total and fractional volumes

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The accurate estimation of cerebellar total volume was made possible by employing cerebellar weight and transforming it into a volume, and by applying the Cavalieri's estimator, taking therefore into account tissue shrinkage. Cerebellar shrinkage secondary to histological processing allows to obtain unbiased stereological estimations insensitive to

- processing-dependent tissue deformations (Dorph-Petersen *et al.*, 2001).
- The estimation of total volume starting from the weight of the
- 191 cerebellum, was performed using the following formula:
- 192 V (cerebellum) = W (cerebellum) / ρ ,
- where ρ refers to the weight-to-volume ratio of cerebellar tissue.
- 194 The estimation of the total volume of the cerebellum through use of the
- 195 Cavalieri principle was conducted by using the test point system (Fig. 2)
- and following the equation below (Howard and Reed, 1998):
- 197 $V = \Sigma P \cdot SSF \cdot T \cdot (a/p)$
- where ΣP is the total number of points hitting the structure; SSF (1/25)
- represents the section sampling fraction; T (25 μ m) is the section thickness
- and a/p $(465,267 \mu m^2)$ refers to the area per point.
- The fractional volume (V_v) of cerebellar structures including white
- 202 matter, grey matter, molecular and granular layers, was estimated using the
- following formula (Gundersen *et al.*, 1988a):
- 204 V_v (structure)= $\sum P$ (structure) / $\sum P$ (cerebellum)
- where $\sum P$ (structure) is the number of points hitting the white matter, gray
- 206 matter, molecular and granular layers, and $\sum P$ (cerebellum) is the number
- of points hitting the cerebellum.
- Lastly, in order to estimate the volume accounted for by each
- structure, each volume fraction was multiplied by the total volume of the

210 cerebellum.

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Estimation of surface area

- The surface density (S_v) of the cerebellum was estimated by using
- 213 test lines (Fig. 2b), and by employing the following formula (Howard and
- 214 Reed, 1998):
- 215 $S_v = 2 \cdot \sum l / (\sum P \cdot l/p)$
- Where $\sum l$ represents the total number of intersections between the outer
- surface of the cerebellum and the test lines, $\sum P$ refers to the points hitting
- 218 the molecular layer, 1/p (658 µm) was the length of each test line associated
- 219 to each point of the test grid.
- 220 Consequently, for estimating the surface area, surface density was multiped
- by the volume of the molecular layer.
- 222 In addition, the thickness (T) of the molecular and granular layers was
- 223 calculated using the following formula (Andersen et al., 2012):
- 224 T (layer) = V (layer) / S (layer)
- where V is the volume and S is the surface area of each layer.

Estimation of Purkinje cell total count

- The optical disector method was employed for the estimation of the
- Purkinje cell total number, and a motorized stage designed by Department
- of Anatomy, Faculty of Veterinary Medicine, of the University of Tehran,
- 230 Tehran, Iran, was employed for the purpose. The microscopic fields were

selected by moving the microscope stage in the x and y directions for a constant distance spanning the entire section thickness. The unbiased counting frame principle was applied for counting the cells. The Purkinje cells whose nucleolus was located inside the counting frame or crossed the accepted lines were sampled, and those whose nucleolus came into focus within disector height were counted (Fig. 3).

- The numerical density of Purkinje cells was calculated using the following formula (Kristiansen and Nyengaard, 2012):
- 239 N_v (Purkinje cells) = $[\Sigma Q^{-}/(a/f \times \Sigma P \times h)] \times t/BA$

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- where ΣQ^{-} represents the total count of Purkinje cells, a/f (9895 μm^{2}) is the area per frame, ΣP is the total number of frames, h (10 μm) is the disector height, t is the sections mean thickness (18.5 μm), measured for each microscopic field, and BA (25 μm) is the block advance.
 - Finally, for the estimation of the total number of Purkinje cells, the numerical density was multiplied by the total volume of the cerebellum, estimated using the Cavalieri's principle.

Estimation of mean Purkinje cellular and nuclear volumes

To estimate the volumes of Purkinje cells and Purkinje cell nuclei, the nucleator technique was utilized (Gundersen *et al.*, 1988b). The volume of the sampled cells was measured by using the unbiased counting frame,

- and following the formula (Gundersen *et al.*, 1988b):
- 252 $V_n = 4\pi/3 \cdot l_n^3$
- Where l_n refers to the intercept length from the nucleolus to the border of
- 254 the cytoplasm (for cellular volume), or to the border of the nucleus (for
- 255 nuclear volume) of Purkinje cells.

Estimation of the coefficient of error (CE)

The precision of the volume estimates, expressed in terms of CE, is related to the variability associated with systematic uniform random sampling (SURS) sampling and point counting of the estimator. The CE for the estimate of the volume (Gundersen and Jensen, 1987), surface area (Kroustrup and Gundersen, 1983) and Purkinje cell count (Braendgaard et al., 1990) was calculated.

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Statistical analysis

All data are expressed as mean ± standard deviation (SD). As for right-skewed distributions, a logarithmic scale was used for individual estimates of Purkinje cellular and nuclear volumes (Weber *et al.*, 1997).

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Results

All cerebella evaluated appeared normal both macroscopically and on histological examination, with all the microscopical structures being distinctly identifiable and without any evidence of pathological processes.

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273 The mean (\pm SD) weight of a guinea-pig cerebellar hemisphere was 274 0.285 ± 0.028 g. The mean volume of a guinea pig cerebellar hemisphere, calculated by dividing the cerebellar weight by its specific gravity, was 275 0.274 ± 0.027 cm³, while the value obtained by employing the Cavalieri's 276 estimator, was 0.110 ± 0.015 cm³. A $61.34 \pm 5.39\%$ total cerebellar volume 277 shrinkage, secondary to the process of paraffin embedding, was estimated. 278 The relative volume fractions of the grey and white matters, expressed as a 279 percentage of total cerebellar volume, were found to be $78.06 \pm 2.66\%$ and 280 281 $21.92 \pm 2.67\%$, respectively. Their absolute volumes, on the other hand, were calculated to be 0.21 ± 0.02 cm³ for the grey matter, and $0.060 \pm$ 282 283 0.006 cm³ for the white matter. The separate and mean values for the above-mentioned parameters, are outlined in Table 1. 284

The surface area of the cerebellum measured $611.4 \pm 96.8 \text{ mm}^2$. The volume of the molecular layer was estimated to be $112.41 \pm 20.56 \text{ mm}^3$ while that of the granular layer $104.38 \pm 7.31 \text{ mm}^3$; the molecular and granular layers accounted for $40.67 \pm 3.87 \%$ and $37.38 \pm 1.77 \%$ of total cerebellar volume, respectively. The mean thickness of the molecular and granular layers was $0.184 \pm 0.020 \text{ mm}$ and $0.169 \pm 0.017 \text{ mm}$, respectively. In Table 2 are shown the mean and individual data calculated for the abovementioned criteria in the six guinea pigs.

The frequency distribution of the Purkinje cellular and nuclear volumes is plotted in Fig. 4. The Purkinje cell volumes were found to be ranging from 987 to 8246.8 μ m³, of which cells with a volume of 3210.1 μ m³ had a higher (13.71%) incidence of occurrence. The estimated volume of Purkinje nuclei was found to be ranging between <117 and 1623.4 μ m³, and nuclei with a volume of 470.9 μ m³ were the most frequently occurring ones (13.54%).

The mean total number of Purkinje cells for a cerebellar hemisphere was calculated to be $253,090 \pm 34,754$ (Table 3).

The mean coefficient of error (CE) and coefficient of variation (CV), along with their ratio (CE²/CV²), calculated for total cerebellar volume, grey and white matter volume fractions, granular and molecular layers volume fractions, cerebellar surface area, and total number of Purkinje cells are shown in Table 4.

Discussion

The mean total volume of a guinea-pig cerebellar hemisphere estimated in the present study is consistent with that calculated in a previous work, which investigated the brain morphology of domestic guinea pigs through quantitative cytoarchitectonic measurements (Kruska, 2014). Cerebellar total volume has been previously assessed by

stereological techniques in other species such as humans, which exhibited a difference between sexes, with male cerebella measuring 120.5 ± 11.1 cm³ in volume, while females 105.9 ± 11.2 cm³ (Taman *et al.*, 2020). Cerebellar volume has also been stereologically estimated in rabbits (Karabekir et al., 2014) and rats (Noorafshan et al., 2018), presenting volumes of 0.69 ± 0.03 cm³, and 0.080 ± 0.004 cm³ for each cerebellar hemisphere, respectively, but also in cats (Sadeghinezhad et al., 2020), presenting a mean cerebellar hemisphere volume of 2.06 ± 0.29 cm³. When comparing total cerebellar volume (in cm³) in relation to body weight (in kg) in each species, it appears that the guinea pigs of the present study have a cerebellar volume to body weight ratio of 0.9, which is consistent with the 0.8 calculated for the rat (Noorafshan et al., 2018), but greater than the 0.4 estimated for the rabbit (Karabekir et al., 2014), and less than an approximate 1.7 for an adult individual of average weight (Taman et al., 2020) and than the approximate 1.1 calculated for a medium-sized cat (Sadeghinezhad et al., 2020).

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The cerebellar weight to body weight ratio was 0.1 in the guinea pig study population, which is in line with an approximate 0.13 calculated for a medium-sized cat (Sadeghinezhad *et al.*, 2020). Cerebellar volumetric modifications have been correlated with physiological factors such as age, gender (Raz *et al.*, 1998), cognitive capability, but also with several

pathological neurological conditions such as Alzheimer's disease, schizophrenia and epilepsy in humans (Bottmer *et al.*, 2005; Sato *et al.*, 2007; Bas *et al.*, 2009; Andersen *et al.*, 2012). A study carried out on rats has also identified a correlation between maternal diabetes and a reduction of total cerebellar volume and thickness of all layers in the offspring (Hami *et al.*, 2016). Volumetric prediction of the cerebellum can therefore find a valuable use in further research on veterinary neurological disease affecting cognition.

The cerebellar gray and white matter volumes have also been sterologically estimated in other species. The total gray matter volume of human cerebella has been calculated to be 88.5 cm^3 , while that of the white matter 22.5 cm^3 (Andersen *et al.*, 2012). Cerebellar grey and white matter volumes were estimated to be $1.46 \pm 0.24 \text{ cm}^3$ and $0.60 \pm 0.06 \text{ cm}^3$, respectively, for the cat (Sadeghinezhad *et al.*, 2020). When compared to the guinea pig and cat, the proportionally more voluminous grey matter in humans can be likely ascribed to their increased development of motor control, coordination, as well as cognitive functions. Moreover, it was noted that, in the early domesticated mammals such as the guinea pig, a decrease in total brain size, which is proportional to the level of encephalization of the species, along with a decrease in total cortex and areas responsible for processing sensory information and motor control,

such as the grey matter, occurred as a consequence of the domestication process, with, however, the cognitive functions not being affected by this change (Kruska, 2005; Kaiser *et al.*, 2015; Welniak-Kaminska *et al.*, 2019). The volumes of the grey and white matter calculated in the present study are markedly greater than those reported by Mallard *et al.* (2000) for neonatal guinea pigs, which is likely due to the large age and body weight discrepancy. The influence of the physiological process of aging on volumetric changes in the cerebellar gray and especially the white matter has been assessed in several studies (Jernigan *et al.*, 2001; Walhovd *et al.*, 2005).

Several human neurological diseases affecting cognition have also been

Several human neurological diseases affecting cognition have also been observed to cause volume losses of the cerebellar gray and white matters (Fennema-Notestine *et al.*, 2004; Anderson *et al.*, 2009), as evidence of the role that the cerebellum plays in cognition.

The mean volumes of the molecular and granular layers in the guinea pig cerebellum estimated in the present work are significantly greater than the corresponding values reported by Mallard *et al.* (2000) for neonatal guinea pigs, and, comparing the two studies, the volumes of the two layers are apparently not proportionally related to body weight. The mean corresponding volumes referring to humans are 54.4 cm³ for the molecular layer, and 37.9 cm³ for the granular layer (Andersen, 2004). The mean

volume of the molecular and granular layers of the cerebellum of normal rats was reported to be $0.035~\rm cm^3$ and $0.024~\rm cm^3$, respectively (Dortaj *et al.*, 2018). In cats' cerebella, the mean molecular layer volume had been reported to be $0.89\pm0.16~\rm cm^3$, while that of the granular layer $0.56\pm0.10~\rm cm^3$ (Sadeghinezhad *et al.*, 2020). The relative proportions of the molecular and granular layers of the cerebellum in the different species seem to be conserved, thanks to the similar cerebellar microscopical anatomy. As a matter of fact, the histological examination of the guinea pig cerebella permitted the clear identification of the molecular, Purkinje and granular layers with their characteristic cellular populations. The conserved volumetric trend seems to be therefore related to function.

A stereological study carried out on intrauterine growth-restricted guinea pigs secondary to placental insufficiency in the second half period of pregnancy, has been seen to cause a reduction in the volume of the molecular and granular layers, as well as in that of the white matter in prenatal guinea-pig cerebella, therefore causing cognitive, motor and behavioral deficits in the post-natal life (Mallard *et al.*, 2000).

When analyzing the distribution of the thickness of the molecular and granular layers in the different subjects comprising our study population, it appears that the measurements are quite consistent and regular, in contrast with what Sultan and Braitenberg (1993) had reported for smaller mammalian species. Andersen (2004) calculated a mean thickness of the molecular layer of $590.00 \pm 0.08~\mu m$ and $410.00 \pm 0.15~\mu m$ for the granular layer in human cerebella. Sadeghinezhad *et al.* (2020), on the other hand, calculated $133.5 \pm 10.1~\mu m$ for the molecular layer and 84.7 $\pm 17.3~\mu m$ for the granular layer in cats' cerebella. Consistently with human and cats' cerebella, the molecular layer appears thicker than the granular, although not in a statistically significant manner; however, it seems that the thickness in guinea pigs is more uniformly-distributed between the two layers when compared to cats and humans' data. This can be explained by different physiological factors such as age. Indeed, a study carried out on cats' cerebella showed that aging causes an increase in granular layer thickness at the expense of that of the molecular layer (Zhang *et al.*, 2006).

With regard to the measurement of the cerebellar surface area, the ratio of cerebellar surface area to cerebellar weight in the different animals comprising the study population remains fairly constant, supporting the proportionality between cerebellar area and cerebellar weight hypothesized by Sultan and Braitenberg for larger mammals (1993), which is probably due, unlike other smaller mammalian species, to the equally constant distribution of grey matter thickness values in our guinea pig population. Further studies on larger population samples are needed to confirm this

finding. The average surface area of the human cerebellum has been previously estimated by different authors to be 550 cm² (Henery and Mayhew, 1989), 1027 cm² (Andersen et al., 2012) and 1160 cm² (Andersen et al., 1992). The human cerebellum, during evolution, underwent a significant expansion of its surface area both in absolute terms as well as in relation to the neocortex; this growth played a critical role in human cognitive development in comparison with other animals, given the role of the cerebellum in cognition (Barton and Venditti, 2014). In the animal kingdom, therefore, it is likely that the cerebellar surface area of highly encephalized species such as higher primates might show a greater development in comparison with mammals of a similar size. On the other hand, a mild but significant reduction in the total cerebellar area has been described in humans with advancing age, showing varying decline trends in the different vermian lobules (Raz et al., 1998). A study carried out on experimentally vitamin C-deprived guinea pig fetuses has revealed a significant reduction in cerebellar surface area due to the obliteration of fissures and the fusion of opposing folia, resulting in a macroscopicallyvisible cerebellar dysplasia in terms of flattening of its surface, analogously to that observed in Lyssencephaly Type 2 (Čapo et al., 2015). The mentioned study is of clinical relevance in pet guinea pigs due to their natural incapability of endogenous vitamin C synthesis (Nishikimi et al.,

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1992), analogously to humans, resulting in the necessity of its dietary supplementation, with the risk of developing vascular as well as neurological disease in case of deprivation.

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Purkinje cells with a perikaryon volume of 3210.1 µm³ and with a 442 nuclear volume of 470.9 μm^3 were found to have the highest frequency of 443 occurrence in the guinea pig cerebellum. Mean Purkinje cellular perikaryon 444 volumes had been estimated to be 12400 µm³ in humans (Korbo and 445 Andersen, 1995), 4900 μm³ (Korbo and Andersen, 1995) and 5600 μm³ 446 (Sørensen et al., 2000) in rats, 17600 µm³ in adult minipigs (Jelsing et al., 447 2006), 2207 µm³ in rabbits (Akosman et al., 2011), and 6994 µm³ in cats 448 (Sadeghinezhad et al., 2020). If considering a mean weight for an adult 449 450 individual of each species, and calculating a ratio of Purkinje volume to 451 body weight, these findings suggest a non-allometric correlation. Indeed, the mini-pig (Jelsing et al., 2006) has a Purkinje volume to body weight 452 ratio that is six times greater than that of humans (Korbo and Andersen, 453 1995), whereas rodents such as the rat (Sørensen et al., 2000) and the 454 guinea pig have, respectively, ratios that are 40 and 180 times 455 proportionally greater than that of humans. The variability encountered 456 457 might be explained by the different degrees of tissue shrinkage (Andersen et al., 1992), by the immersion time of the tissue in the fixative (Jelsing et 458 al., 2006), by the degree of postnatal development of Purkinje perikaryon 459

volume (Jelsing *et al.*, 2006), or by different degrees of significance of Purkinje cell roles in motor, sensory and cognitive functions among the different species.

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The mean total number of Purkinje cells calculated in the present work is consistent with the value reported for the whole cerebellum in a previous work carried out on neonatal guinea pigs, that is in the order of 500,000 (Mallard et al., 2000). It has been demonstrated that the brain of newborn guinea pigs, species characterized by its precocity, presents a high degree of neurological maturity, and that postnatal cerebellar neurogenesis is minimal (Altman and Das, 1967). As a matter of fact, all cerebellar layers, including Purkinje cells, as well as white and gray matters, are well developed and differentiated as early as 45 days post conception (Silva et al., 2016), and that all cerebellar cell proliferation events are entirely complete at birth in this species, unlike other similar rodents (Lossi et al., 1997). In the adult mini-pig cerebellum, on the other hand, the total number of Purkinje cells was in the order of 2.8 million (Jelsing et al., 2006). The numerosity of the Purkinje cell count in the above-mentioned study was, indeed, partially explained by a significant postnatal development in total Purkinje cell number and perikaryon volume, as it had also been demonstrated in rats (Altman and Bayer, 1978), humans (Miyata et al., 1999), and cats (Vastagh et al., 2005). The total number of Purkinje cells in

the whole adult rat cerebellum was estimated at around 320,000 cells (Sonmez *et al.*, 2010), which is markedly less than the value obtained for the guinea pig, and that could be explained by the complex heterogeneity of guinea pigs' Purkinje cells. It has been noted that Purkinje cells in the guinea pig cerebellum show a complex expression pattern of zebrin II, an immunohistochemical marker of cerebellar compartmental heterogeneity, showing three levels of zebrin II expression (Larouche *et al.*, 2003), as opposed to rats, where zebrin II expression only distinguishes two classes of Purkinje cells (Brochu *et al.*, 1990).

The hypothesis that less voluminous brains tend to have a higher cellular density than larger brains (Mwamengele *et al.*, 1993) does not seem to always be applicable, as is the case with the higher count of Purkinje cells in the guinea pigs comprising the present study when compared with the values reported for the rat (Sonmez *et al.*, 2010). Reports of acquired cerebellar degenerative disease in pet guinea pigs, mostly secondary to an infectious etiology, have been described in the literature, with ataxia and loss of voluntary motor control being common clinical signs, and meningoencephalitis and cerebellar cortical hypoplasia with necrosis of granule and Purkinje cells the principal histopathological findings (Monjan *et al.*, 1971; Van Andel *et al.*, 1995; Brabb *et al.*, 2012; Gentz and Carpenter, 2012; Hawkins and Bishop, 2012).

In conclusion, the present study represents the first detailed description of the morphometrical features of the guinea pig cerebellum using design-based stereological techniques. The reference morphometrical data provided for cerebellar structures might find a use as basic anatomical contribution to a greater understanding of neurological diseases when examining cerebellar pathology with relation to function in this exotic pet species of increasing veterinary interest. In addition, the present study might prove useful by providing a comparison with available data in humans and other mammals for future research investigating the basis of motor, cognitive and behavioral diseases in the different species.

Conflict of interests

517 The authors have no conflict of interests to declare.

Author contributions

519 M.D.S.: acquisition of data, data analysis/interpretation, drafting of the

manuscript; J.S.: concept/design, acquisition of data, data analysis/interpretation, critical revision and approval of the manuscript; J.R.N.: data analysis/interpretation, critical revision of the manuscript and approval of the article; M.A.A.: data analysis/interpretation; A.S.: data analysis/interpretation; N.D.S.: acquisition of data; R.C.: concept/design; critical revision of the manuscript and approval of the article; A.G.: acquisition of data, critical revision of the manuscript and approval of the article.

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Table 1. Stereological data for total volume of cerebellar hemisphere and proportional volume of gray matter and white matter in six guinea pigs.

Animals	Cerebellu m weight (g)	Total volume of cerebellum (weight/sp ecific gravity) (cm³)	Total volume of cerebellum (Cavalieri estimator) (cm³)	Shrinkage (%)	Gray	matter	White matter	
					Volume fraction (%)	Volume (cm³)	Volume fraction (%)	Volume (cm³)
1	0.257	0.247	0.117	54.47	75.94	0.1875	24.05	0.0594
2	0.282	0.271	0.118	58.15	80.42	0.2179	19.57	0.0530
3	0.298	0.286	0.117	60.73	77.60	0.2219	22.39	0.0640
4	0.263	0.252	0.079	69.96	78.64	0.1981	21.35	0.0538
5	0.280	0.269	0.112	60	74.37	0.2000	25.62	0.0689
6	0.335	0.322	0.118	64.77	81.42	0.2621	18.57	0.0597
Mean±SD	0.285 ± 0.02	0.274 ± 0.02	0.110 ± 0.015	61.34±5.39	78.06 ± 2.66	0.2145 ± 0.02	21.92±2.66	0.0598 ± 0.00
	8	7				66		60

Animals	1	2	3	4	5	6	Mean±SD
Surface area (mm ²)	486.066	555.984	630.003	627.302	592.925	776.140	611.40±96.8
Volume fraction of molecular layer (%)	40.28	42.24	39.3	41.63	34.42	46.19	40.67±3.87
Volume of molecular layer (mm ³)	99.4	114.4	112.3	104.9	92.5	151.0	112.41±20.56
Volume fraction of granular layer (%)	35.65	38.18	38.29	37.01	39.94	35.23	37.38±1.77
Volume of granular layer(mm³)	88.0	103.4	109.5	93.2	107.4	113.4	104.38±7.31
Thickness of molecular layer(mm)	0.204	0.205	0.178	0.167	0.156	0.194	0.184±0.020
Thickness of granular layer(mm)	0.181	0.185	0.173	0.148	0.181	0.146	0.169±0.017

Table 3. Stereological data for numeral density and total number of Purkinje cells in cerebellar hemisphere in six guinea pigs.

Animals	1	2	3	4	5	6	Mean±SD
Numeral density (cells/mm³)	2532	2413	2215	2546	2197	1986	2314.833 ± 220.099
Total number	296010	284380	258570	200660	245280	233640	253090 ± 34754

Table 4. The mean coefficient of error (CE) and coefficient of variation

	Total volume	Grey matter volume	White matter volume	Granular layer volume	Molecular layer volume	Surface area	Total number of Purkinje cells
CE	0.016	0.080	0.050	0.033	0.031	0.0162	0.080
CV	0.140	0.123	0.101	0.096	0.182	0.158	0.137
CE^2/CV^2	0.013	0.423	0.252	0.121	0. 029	0.479	0.346

962 (CV) of stereological analysis of guinea pig cerebellum (n=6)

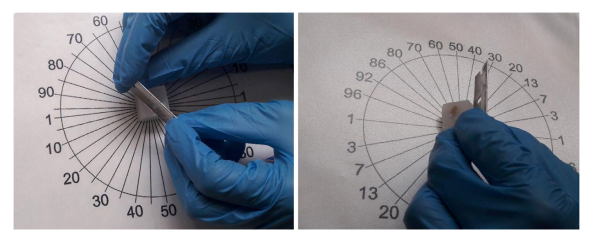


Figure 1. Isotropic, uniform random sections of the guinea-pig cerebellar hemispheres were obtained by applying the orientator method. (a): A randomly chosen cerebellar hemisphere for each animal was embedded in a paraffin block and placed at the centre of a circle with 90 equidistant divisions along the perimeter. A random number between 0 and 90 was looked up and the paraffin medium was cut along a line parallel to the direction of the selected number (here, 75). (b): The block was placed on its cut surface at the center of a second circle, with 96 nonequidistant divisions along its perimeter. The paraffin was cut along a line parallel to the direction of a random number ranging from 0 to 96 (here, 50), and the block was finally re-embedded in paraffin while placed on its cut surface, and consecutive 25 μm-thick sections were cut with a microtome.

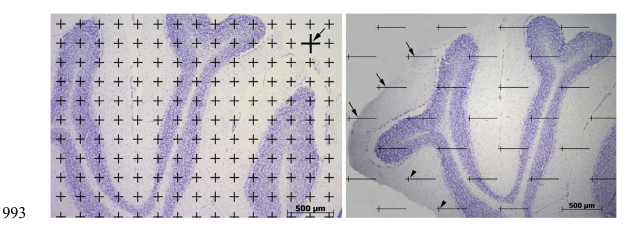


Figure 2. Estimation of cerebellar volume and surface area by employing the point-counting and the test-lines systems. (a): The volume of the cerebellar structures was estimated by randomly superimposing a point-counting probe onto each section. The upper right corner of each point (arrow) was taken as a reference for the count of the number of points hitting the region of interest. (b): The surface area of the cerebellum was estimated by superimposing test-lines onto each section. The arrowheads

show two points hitting the molecular layer, whereas the arrows indicate

the intersection between test lines with the outer cerebellar surface.

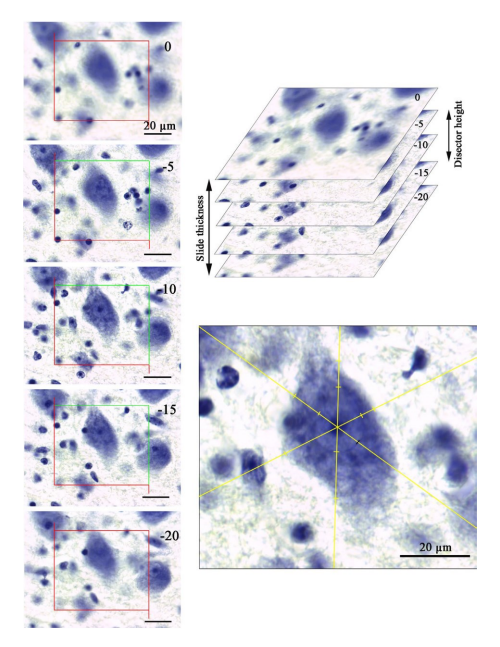


Figure 3. Use of the optical disector technique for the Purkinje cell count and of the nucleator technique for the estimation of the Purkinje cellular and nuclear volumes. (a): The microscopic fields were selected by moving the microscope stage in the x and y directions for a constant distance. Then, the stage of microscope moved in z-axis and the consecutive focal planes were evaluated within optical disector height (10 μ m from -5 to -15 μ m).

(b-f): The unbiased counting frame principle was applied for the Purkinje ell count. The cells whose nucleolus was located inside the counting frame or crossed the accepted lines were sampled, and those whose nucleolus came into focus within disector height were counted. (g): The intercept length from the nucleolus to the border of the cytoplasm, or to the border of the nucleus, was measured for the estimation of Purkinje cellular and nuclear volumes, respectively.

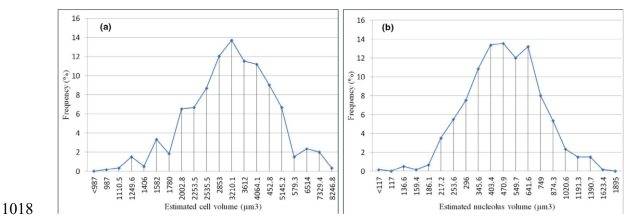


Figure 4. Graphs showing the frequency distribution of the Purkinje cellular (a) and nuclear (b) volumes in the guinea-pig cerebellum.