Novel human melanoma brain metastasis models in athymic nude fox1nu mice: Site-specific metastasis patterns reflecting their clinical origin

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Abstract

Background: Malignant melanomas frequently metastasize to the brain, but metastases in the cerebellum are underrepresented compared with metastases in the cerebrum.

Methods: We established animal models by injecting intracardially in athymic nude fox1nu mice two human melanoma cell lines, originating from a cerebral metastasis (HM19) and a cerebellar metastasis (HM86).

Results: Using magnetic resonance imaging (MRI), metastases were first detected after a mean of 34.5 days. Mean survival time was 59.6 days for the mice in the HM86 group and significantly shorter (43.7 days) for HM19-injected animals (p < 0.001). In the HM86 group, the first detectable metastasis was located in the cerebellum in 15/55 (29%) mice compared with none in the HM19 group (p < 0.001). At sacrifice, cerebellar metastases were found in 34/55 (63%) HM86-injected mice compared with 1/53 (2%) in the HM19-injected (p < 0.001) mice. At that time, all mice in both groups had detectable metastases in the cerebrum. Comparing macroscopic and histologic appearances of the brain metastases with
INTRODUCTION

The human skin is the largest organ in the body and its melanocytes offer protection against radiation from the sun.\textsuperscript{1–3} Regrettfully, this protection is incomplete and sun exposure is the number one primary cause of cutaneous malignant melanoma (MM) in Caucasians.\textsuperscript{4,5} Worldwide, the incidence of MM has increased in Caucasian populations over the last 15 years\textsuperscript{6–10} and it will likely increase further if preventive efforts are not undertaken.\textsuperscript{11} In Norway, the incidence has increased nearly 15-fold since the Norwegian Cancer Registry was established in 1952, and in the age group 15–54 years, MM is now the second most common cancer type for both sexes. In 2015, the age-adjusted incidence rate for men and women was 39.5 and 35.5 per 100,000 person-years, respectively,\textsuperscript{12} making Norway one of the countries with the highest incidence of this malignancy.\textsuperscript{13}

Brain metastases can occur with most cancer types.\textsuperscript{14} In clinical series, 80\% of the brain metastases are found in the cerebrum and 20\% in the posterior fossa (cerebellum and brain stem).\textsuperscript{15} This distribution closely parallels the regional blood flows and blood volumes as well as the relative brain tissue mass of the respective regions.\textsuperscript{16,17} Cutaneous MM frequently metastasize to the brain\textsuperscript{22} in contrast to uveal melanomas.\textsuperscript{23} Interestingly, Rogne et al.\textsuperscript{14} found that brain metastases from MMs were significantly underrepresented in the cerebellum, accounting for only 5\% of all the intracranial MM metastases resected in their neurosurgical series, and not 15\%–20\% as might be expected. This reduced propensity of MM to establish growth in the cerebellum is likely associated with microenvironmental and tumor cell-associated factors,\textsuperscript{14,24} in accordance with the seed and soil theory of metastasis. However, it is difficult to study associations between molecular and immunologic tumor characteristics and the occurrence and biologic behavior of brain metastasis in clinical series.\textsuperscript{18}

Preclinical studies of brain metastases have been limited by the scarcity of clinically relevant models.\textsuperscript{19} We have previously reported several brain metastasis models in immunodeficient mice and rats\textsuperscript{20,21}; however, none of these are suited for studying tissue-specific growth within the brain. We have established two in vitro cell lines directly from human MM brain metastases, of which one originates from a cerebellar and one from a cerebral metastasis in the respective patients. Melanoma brain metastasis animal models were developed by injecting the cell lines intracardially in female athymic nude fox\textsuperscript{1}\textsuperscript{nu} mice. Our first objective was to examine whether the pattern of spread observed in the clinic could be reproduced in mice. The appearance and growth of resulting metastases in the animals were followed by MRI, examining patterns of metastasis to the cerebrum and cerebellum. The second objective was to study the relationship between tumor phenotype and metastatic site by morphologic and immunohistochemical analyses, comparing the mouse brain metastases to each other and to the clinical metastases from which the cell lines originated.

METHODS

2.1 | Human tissue collection and mutation analyses

Fresh biopsy material was obtained during surgery at the Department of Neurosurgery, Oslo University Hospital. The tumor tissue was collected in 50 ml tubes filled with RPMI 1640 (Merck KgaA, Sigma-Aldrich, catalog number R0883) and transported to the Department of Tumor Biology, Institute for Cancer Research, Oslo University Hospital. All patients had signed a written consent before tumor material was collected and the study was approved by The Regional Ethical Committee, Region South-East,
Norway (approval number 2011/2183). Immediately upon arrival in the laboratory, the tissue was split into samples for subsequent molecular analyses, which were stored at \(-70^\circ\text{C}\), samples for evaluation of tumor cell content, and one sample for the establishment of cell lines. DNA and RNA were isolated using the AllPrep DNA/RNA/miRNA Universal Kit (Qiagen GmbH, catalog number 80224), and the QiaCube system. Targeted DNA sequencing was conducted using the Ion AmpliSeq Cancer Hotspot Panel (v2) (LifeTechnologies, catalog number 4475346) covering \(~2800\) hotspot mutations in 50 cancer-related genes. Data from the Ion PGM sequencing platform runs were processed by The Torrent Suite Variant Caller using panel customized parameters. Additional cutoff filters (e.g. minimum allele frequency) and manual evaluation by visual inspection were applied. In the patient brain metastases from where the two cell lines were established, the following gene aberrations were detected: HM19– BRAF K601E, TP53 S241F, APC R1105R, VHL P138S, HM86– NRAS Q61K.

2.2 | Cell culture

Cell lines from two different patients were developed; HM19 originated from an MM metastasis in the cerebrum, whereas HM86 was derived from an MM metastasis in the cerebellum. Approximately one-third of the tissue received was chopped into small pieces with a scalpel in a petri dish. The small tissue pieces were resuspended in the transport medium (RPMI 1640) and filtered through a 70 µm Falcon filter (Corning Incorporated, catalog number 431751) into a new 50 ml tube. The petri dish was washed with 2–5 ml RPMI 1640 supplemented with 10% FBS (Sigma-Aldrich, catalog number F7524). The filter was also washed with this solution and all collected in the 50 ml tube. After centrifugation at 235 g for 5 min, the pellet was resuspended in a 5 ml cell culture medium and placed in a 25 cm² cell culture flask (Nunc, Thermo Fisher Scientific, catalog number 156367). The cells were cultivated in RPMI supplemented with 10% FBS and 2 mM stable Glutamin (Merck KgaA, Sigma-Aldrich, catalog number G8541) at 37°C with 5% CO₂. The medium was changed once a week. After some weeks in culture, and when appropriate confluence was achieved, the cells were detached by 0.01 M EDTA (Merck KgaA, Sigma-Aldrich, catalog number E8008) and split into new flasks. After 10 passages, we consider a cell line to be established. Both cell lines were routinely tested for Mycoplasma (VenorGem Mycoplasma detection kit, Minerva Biolabs GmbH, catalog number 11-1100), while the human origin and identity were confirmed by genomic fingerprinting using the PowerPlex 16 kit (Promega, catalog number DC6530). For LV injections in mice, the cells were detached by EDTA, washed once, and resuspended in RPMI without FBS to a concentration of \(5 \times 10^5\) cells/0.2 ml.

2.3 | Animals

In our nude rodent facility at the Department for Comparative Medicine, we bred female athymic nude fox1nu mice \((N = 120,\) aged \(>4\) weeks). No more than 10 animals were placed in each cage. To minimize infection, the cages were kept in a pathogen-free environment at constant temperature (21.5 ± 0.5°C) and humidity (55 ± 5%), 15 air changes/h, and a 12 h light/dark cycle. All procedures and experiments involving animals were approved by The National Animal Research Authority (FOTS ID: 7310) and carried out according to the European Convention for the Protection of Vertebrates used for Scientific Purposes. Food and water were daily controlled.

2.4 | Left ventricular injection of the heart

The mice were anesthetized with a subcutaneous injection of Zoletil* 5 µg/g body weight (Virbac Laboratories, catalog number A5727) and placed in a supine position on a styrofoam board. Parasternal on the left side, an incision was made to localize the third intercostal space where the punction site is. The tumor cells were introduced to the left ventricle of the heart by a 27-gauge cannula connected to a 10 cm plastic tube (Valu-set, BD Bioscience, Franklin Lakes, catalog number 387412). This arrangement was mounted on a stereotactic instrument to increase the accuracy. Because of the increased pressure in the left ventricle, we could assess if the punction was successful by observing oxygenated blood in the plastic tube. Tumor cells \((5 \times 10^5)\) in 200 µl were injected. After the injection,
we sealed the skin with one absorbable suture (Polysorb 5-0; Covidien, catalog number GL-890) and Histoacryl® (B. Braun, Rubi, Spain, catalog number 1050052), after observing pulsating blood in the plastic tube. After the 10 min procedure, the animals were placed in cages often.

2.5 | Magnetic resonance imaging

MRI was carried out at the MRI Core Facility at Department for Comparative Medicine, Oslo University Hospital, by using a Bruker Biospec 7 Tesla small-animal scanner. An anatomically shaped receive-only coil array (2 × 2) was placed over the head of the animal (Bruker BioSpin), and an 86 mm 1H volume resonator was used as a transmit coil. Mice were sedated with 5% Sevofluran® (Baxter, catalog number 460-220-12) in O2 and maintained under general anesthesia (3% Sevofluran®). Respiration rate and body core temperature were monitored throughout the procedure by using an abdominal pressure-sensitive probe and a rectal temperature probe (Small Animal Instruments).

Coronal and axial T2-weighted images were obtained by using a fast spin-echo sequence (RARE) with a repetition time (TR) of 2500 ms, an echo time (TE) of 35 ms, a RARE factor of 8, an image matrix of 256 × 256, a field of view (FOV) of 2 × 2 cm², a slice thickness of 0.5 mm, a slice gap of 0.3 mm, and averages of 2. Scan time was 2:40 min.

MRI was used to locate the sites of metastatic intracranial lesions and to monitor the tumor growth. An initial baseline MRI scan was performed 24 h after injection of tumor cells to evaluate normal brain anatomy. The animals were then scanned once weekly for 5 weeks, and thereafter every tenth day until sacrifice.

The following additional parameters were recorded: location of the first intracranial lesion, time from intracardiac injection until first intracranial metastasis, locations and number of subsequent metastases, and time from injection until sacrifice.

The experimental endpoint was determined by the onset of neurologic symptoms, unacceptable weight loss (>15%) and body condition, or hind-limb paralysis. At the endpoint, the mice were again scanned and subsequently sacrificed by cervical dislocation.

2.6 | Animal tissue collection and preparation

After the experimental endpoint, the brain was surgically removed from the cranium. We made an incision in the level of C3 and dissected our way to the foramen magnum. From the foramen magnum, we performed a craniectomy from the medulla oblongata to the olfactory bulb. By stump dissection, the brain was free and placed on a sterile glass slide, where it was photographed and immediately placed in a solution of formalin and sent to the Department of Pathology, Oslo University Hospital for 3 days for sectioning and immunohistochemical analyses.

For the immunohistochemical staining, the Dako EnVision FLEX system (DakoCytomation, catalog number GV82311-2) was used. Formalin-fixed paraffin-embedded sections (thickness 3 μm) from mouse brains were treated (97°C for 20 min and cooled down to 65°C) with the PT-link and FLEX Target Retrieval Solution (High pH = 9.0)
for deparaffinization, rehydration, and target retrieval (catalog number GV80411-2). To block endogenous peroxidase, the sections were treated with EnVision Peroxidase-Blocking Reagent (catalog number S202386-2) for 5 min. The sections were incubated either with Melan-A antibody (Abcam, catalog number GR111252-2) dilution 1:250, S100 antibody (DakoCytomation, catalog number #27862) dilution 1:1500 or with anti-HMB-45 (DakoCytomation, catalog number #4684) dilution 1:75 for 30 min.

The sections were then incubated for 30 min with either labeled Polymer HRP antimouse (Dako) for HMB-45 or labeled Polymer HRP antirabbit (Dako) for Melan-A and S100. Then they were stained for 10 min with 3′3′-diaminobenzidine tetrahydrochloride (DAB), counterstained with hematoxylin, dehydrated, and mounted in Cytoseal XYL (Thermo Fisher Scientific, catalog number 8312-4). The sections were deparaffinized for 10 min with xylol followed by rehydration for hematoxylin and eosin (HE) staining. Subsequently, the sections were stained for 6 min with hematoxylin and then cleaned with water before they were stained for 1 min with eosin. Diatex was used for dehydration and mounting the sections.

Sections were analyzed using a Mirax Scanner, Zeiss Microsystems GmbH. Scanning objective 20×, resolution 230 nm/pixel.

2.7 | Data analysis

All images were analyzed using the open-source DICOM viewer OsiriX (OsiriX version 7.0.2). Scans taken at the endpoint were examined retrospectively after sacrifice, allowing for easier identification of the seminal lesion. MRI scans were controlled by a senior neurosurgeon.

2.8 | Statistics

Statistical analyses were performed using SPSS 21 (IBM). A sample size calculation was done in order to evaluate how many animals were needed in each group in order to get significant results. We performed a chi-squared test with the following parameters; $\alpha = 0.05$ and $\beta = 0.2$ (power $= 1 - \beta$) to calculate the needed sample size. According to the power analysis, 49 mice per cell line were needed. Allowing for 20% incidental loss of animals, the total sample size was 120 mice, resulting in 60 mice in each group for metastasis evaluation. Nine mice died after the injections and five mice failed to develop metastases in the HM 86 group, and seven in the HM 19 group, leaving 108 mice included in this study. We used the chi-square test when comparing the distribution of metastases between the two groups. Survival and time to first detected metastases were measured using a Kaplan–Meier plot. A log-rank test was used to evaluate the $p$-value.

3 | RESULTS

One hundred and twenty female athymic nude fox1 nu mice were injected, of which nine died after intracardiac injections. Nine new animals were then injected, for a total number of 120. Twelve of 120 animals failed to develop any metastases (10%), leaving 108 mice for further analysis. The mean time from cell injection to first MRI-detected intracranial metastasis was 34.5 days. By using a log-rank test, we found no statistically significant difference between the two groups injected with the different cell lines ($z = 2.03$, $p = 0.423$) (Figure 1).
intracranial metastasis was in the cerebrum. The lesions were well demarcated and widely distributed within the brain. Within weeks 4–8, the lesions did grow in both size and number. Animals in the HM19 group had a consistent and increasing development of symptoms with time. After an average of 43.8 days, the mice had either developed hind-limb paralysis, respiratory failure, or a weight loss >15% and were sacrificed (Figure 2). At the time of sacrifice, only one mouse (2%) had developed metastasis in the cerebellum (Figure 3). Notably, 15 animals (30%) injected with HM19 cells eventually developed leptomeningeal lesions. Animals with these types of lesions typically expressed elevated respiratory frequency, cyanosis, and fatigue presumably due to respiratory failure secondary to brain stem compression.

Fifty-five mice injected with HM86 cells were available for analysis. In this group, the first detected intracranial metastasis was in the cerebellum of 15 mice (29%). From week 4 to week 6, the lesions did increase slowly in size and number, but from week 7 onward, the lesions did grow rapidly and the animals in this group had a dramatic and rapid course of their illness. The mean overall survival time in this group was 59.6 days, significantly longer than with HM19 cells (log-rank test: $z = 4.84$, $p < 0.001$) (Figure 2). At the time of sacrifice, as many as 34 animals (62.9%), had developed cerebellar metastases, but at that time, all mice also had lesions in the cerebrum (Figure 3).

Using the chi-square test, we found a significant relationship between the distribution of metastases in the cerebellum between the two cell lines. HM86 are more likely than HM19 to metastasize to the cerebellum, $X^2 (1, N = 108) = 44.26$, $p < 0.01$.

### 3.2 Metastasis morphology and immunohistochemistry

Macroscopic examination of the mouse brains confirmed the location and size of the metastases detected by MRI, as exemplified in Figure 4. H&E sections from the macroscopic metastases in mice were taken in a plane similar to the corresponding MRIs to make them comparable to the imaged tumors (Figure 5). Tissue samples from the original brain metastases in the patients as well as brain metastases in mice were stained with melanoma markers Melan-A, S100, and HMB-45 and were all dominated by positive cells. Comparing the morphologic appearance in tumor sections (Figures 5 and 6), we found that the metastases arising from cell line HM19 had a nodular pattern with cells growing in a nest-like pattern, with large necrotic areas and mitoses. These features were evident in samples both from the patient and mouse metastases. Metastases arising from the HM86 cell line differed

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**FIGURE 5** The left column represents mice injected with HM19 and the right with HM86. Figures 1 and 2 are $T_2$-weighted MRI scans. Arrows point to the metastases. Pictures A, B, and C show the cerebral metastasis in a mouse injected with HM19 cells at increasing magnification and standard H&E staining (scale bar: A: 1000 µm, B: 200 µm, C: 100 µm) and picture D shows an IHC analysis of the same metastasis stained for S100 (Scale bar 100µm). Picture E and F show a cerebral metastasis in a mouse injected with HM86 cells at increasing magnification (scale bar: E: 1000 µm, F 200 (scale bar E: 1000 µm, F: 200 µm). There is a distinct difference in growth pattern compared with the corresponding cerebellar metastasis (G) (scale bar: 200 µm). Picture H shows the same lesion as depicted in G, but at higher magnification (scale bar: 50 µm), and stained for S100
from HM19 and showed a more epithelioid growth pattern as well as cells with spindle-like features. The latter was most prominent in the cerebral lesions, whereas the cerebellar lesions showed a more typical epithelioid picture. These tumors also had fewer necrotic areas, but the same amount of mitoses. The HM86 cells were less...
pigmented than the HM19 cells in both the human material and in metastases in mice.

4 | DISCUSSION

It is well known that different cancer types have different propensities for metastasizing to the brain, but most studies do not differentiate between the locations of the metastasis within the various brain compartments. Rogne et al.\(^1\) reported that in a large clinical series of patients with MMs, cerebellar metastases were significantly underrepresented compared with what is reported for other cancer types. If blood flow or tissue volume was the decisive factor, MM metastases should occur in the cerebrum in 80% of the cases, brain stem in 5%, and cerebellum in 15%. However, Rogne et al.\(^1\) found that the rate of cerebellar MM metastases was as low as 5%. To identify underlying causes of this difference, one has to search among the multitude of factors that may affect the metastatic process to the brain.\(^2\)

We wanted to develop models that allow the study of this biologically interesting phenomenon. For this purpose, we used intraventricular heart injections in athymic nude fox1\(^{nu}\) mice of two melanoma cell lines originating from a cerebral metastasis (HM19) and a cerebellar metastasis (HM86). We postulated that the HM19 cell line would follow a similar pattern in an experimental brain metastasis model in immune-deficient mice, whereas the HM86 cell line would show a higher propensity to metastasize to the cerebellum. We initially used contrast-enhanced MRI (CE-MRI), but the metastases were better visualized in a regular \(T_2\)-weighted sequence (Figure 7). Our observation corresponds with those of several studies that describe a low sensitivity of CE-MRI in animal models because of an intact BBB early in the metastasis development, or where the tumor grows to utilize preexisting capillaries.\(^2\)\(^6\)\(^-\)\(^9\) Serial \(T_2W\) MRI proved excellent to detect and follow the development of intracranial spread in the mice, and two distinct patterns of metastasis for the two MM cell lines were observed, reflecting their origins.

We found a significant overrepresentation of cerebellar metastases in mice injected with the HM86 cell line compared with those injected with HM19 cells using a chi-square test \((X^2 (1, N = 108) = 44.26, p < 0.01)\). This finding is remarkable and warrants further studies of the underlying factors involved. Notably, the metastases in the mice were also morphologically similar to their respective original human brain metastases. The cerebellar tumors were more nodular and less vascularized as well as well demarcated compared with the cerebral ones. Classical melanoma markers like Melan-A and S100 were positive in both cell line tumors and in the tumors of origin.

Interestingly, as many as 28% of the HM19 mice had leptomeningeal metastases at the time of sacrifice (Figure 8). Such metastases may become manifest first at a late stage of the disease, possibly not being easily observed in cases with very aggressive disease. However, it is known that clinical leptomeningeal MM metastases are associated with rapid disease progression with a mean survival of only 8–10 weeks after being diagnosed.\(^3\)\(^0\) The HM19 cell line may provide a missing research tool also for studying the biology of this disease manifestation.\(^1\)\(^7\),\(^1\)\(^9\),\(^3\)\(^0\),\(^3\)\(^1\)

The intracardiac injection of cells results in hematogenous spread throughout the animals, in our case with a very good take rate (90%). However, this approach also has its clear limitations as 500,000 tumor cells were injected over a very short period into the systemic circulation, a situation that can lead to intravascular cancer cell aggregates, and misguided injections may result in perfusion to the pulmonary circulation or thorax. In both cases, this can lead to premature death, which occurred in nine of our mice (7%). Furthermore, the wide distribution of cancer cells may give rise to metastases anywhere in the body, which might lead to death before brain metastases are detectable. An alternative approach to obtain experimental brain metastases is to inject the cells directly into the internal carotid artery.\(^2\)\(^0\),\(^3\)\(^2\) This provides a high tumor load to the brain but may be less suitable to study homing mechanisms. Intracarotid injection also demands microsurgical skills, is more time-consuming, and possesses a higher risk of mortality or stroke. In addition to the two methods mentioned above, intrathecal injections give meningeal tumors,\(^3\)\(^3\),\(^3\)\(^4\) and last, cells may also be injected
orthotopically into the brain.35 Despite the limitation associated with the intracardiac injection model, the approach gives several advantages compared with other available options for studying brain metastases, a notion supported by the present results. Thus, only one of 53 mice injected with HM19 cells, originating from a cerebral metastasis, developed cerebellar metastases in the mice, compared with 34 of 54 animals injected with HM86 cells from a cerebellar metastasis. That all HM86-injected mice also had cerebral manifestations at the end of their somewhat long life span may reflect that over time absolute tissue specificity is very rare.

Microenvironmental factors are essential for metastatic growth and include among other release of soluble factors and bidirectional interaction between cancer cells and the microenvironment.36–38 Such factors may affect tumor cell growth, motility, invasion, and angiogenesis39–41 and might be mediated, for example, by expression of cell surface molecules encoding receptors or ligands,42 microRNAs,43 exosomes,44 and metabolic reprogramming.45,46 The preference of HM86 cells for growth in the cerebellum is likely the result of a complex interplay between tumor cells and microenvironmental factors. Such factors may also explain the small differences in morphology between HM86 metastases in the cerebellum and cerebrum.

In summary, the present work demonstrates that human brain-metastatic melanoma cells, even after being cultured in vitro for a long period, have retained important characteristics in their interaction with subtle micro-environmental brain tissue-associated differences in mice. The two new models offer novel possibilities for investigating tumor- and host-associated factors involved in MM brain metastasis in general and in determining tissue-specific metastasis in particular. This is in line with the comments in a recent review paper in which the authors emphasize the need for new animal models for studying brain metastasis.18

ETHICS STATEMENT
All patients have signed a written consent form before tumor material was collected and the study was approved by the Regional Ethical Committee, Region South-East, Norway (approval number 2011/2183). All procedures and experiments involving animals were approved by the National Animal Research Authority (FOTS ID: 7310) and carried out according to the European Convention for the Protection of Vertebrates used for Scientific Purposes.

ACKNOWLEDGMENTS
We would like to thank the internal research education course for medical students (Forskerlinjen) at the University of Oslo for funding this project. We thank the Department of Pathology at the Oslo University Hospital, especially Mette Forsund, for preparing the histologic materials. Last, but not least, we would like to thank Sindre Lee-Ødegård for helping us with the statistics.

CONFLICT OF INTEREST
No conflict of interest.

DATA AVAILABILITY STATEMENT
The data used to support the findings of this study are included in the article. Raw data are available from the corresponding author, upon reasonable request.

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**How to cite this article:** Svendsen HA, Meling TR, Nygaard V, et al. Novel human melanoma brain metastasis models in athymic nude fox1nu mice: Site-specific metastasis patterns reflecting their clinical origin. *Cancer Med*. 2021;10:8604–8613. doi:10.1002/cam4.4334.