Carbon nanotubes introduced into the abdominal cavity of mice show asbestos-like pathogenicity in a pilot study

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Carbon nanotubes1 have distinctive characteristics2, but their needle-like fibre shape has been compared to asbestos3, raising concerns that widespread use of carbon nanotubes may lead to mesothelioma, cancer of the lining of the lungs caused by exposure to asbestos4. Here we show that exposing the mesothelial lining of the body cavity of mice, as a surrogate for the mesothelial lining of the chest cavity, to long multiwalled carbon nanotubes results in asbestos-like, lengthdependent, pathogenic behaviour. This includes inflammation and the formation of lesions known as granulomas. This is of considerable importance, because research and business communities continue to invest heavily in carbon nanotubes for a wide range of products⁵ under the assumption that they are no more hazardous than graphite. Our results suggest the need for further research and great caution before introducing such products into the market if long-term harm is to be avoided.

Carbon nanotubes (CNTs) are often considered to epitomize the field of nanotechnology—a diverse collection of nanoscale technologies that are projected to be associated with \$2.6 trillion worth of manufactured goods by the year 2014 (ref. 6). The global market for CNTs is predicted to grow to between \$1 billion and \$2 billion by 2014, spurred on by new and increasing industrial demands⁵. Meanwhile, widespread concerns have been raised that a poor understanding of how to safely develop and use engineered nanomaterials—including carbon nanotubes—could undermine business interests and unduly jeopardize human health and the environment⁷.

The unique nanometre-scale structure of CNTs is based on a graphene cylinder, typically a few nanometres in diameter, which can range in length from a few micrometres to millimetres³. Single-walled nanotubes (SWNTs) consist of one such cylinder, and multiwalled nanotubes (MWNTs), as used in this study, comprise 2 to 50 such cylinders concentrically stacked with a common long axis. This structure gives nanotubes an unusual

combination of properties that are highly desirable in many industrial products^{3,8}. Their high aspect ratio (ratio of length and width) makes them an attractive structural material, but their nanometre-scale diameter and needle-like shape have drawn comparisons with asbestos^{9,10}.

Exposure during mining and the industrial use of asbestos led to a global pandemic of lung diseases. Study of disease in exposed populations showed that the main body of the lung was a target for asbestos fibres, resulting in both lung cancer and scarring of the lungs (asbestosis). The outside surface lining of the lung and its associated tissue, the pleura, was found also to be a target, with cancer of the pleura (mesothelioma), fluid accumulation in the pleural space (effusion) and scarring of the pleura (pleural thickening and plaque formation) being found in association with asbestos exposure11. A critical factor underlying this pandemic is a prolonged latency period between exposure and the development of mesothelioma, the hallmark cancer of asbestos exposure. Toxicologists have derived a paradigm in which a hazardous fibre is one that is thinner than 3 µm, longer than ~20 µm and biopersistent in the lungs, in other words not dissolving or breaking into shorter fibres12. Above all, for there to be any adverse effect, the numbers of such fibres must reach a sufficient level to cause chronic activation of inflammatory cells. genotoxicity, fibrosis and cancer in the target tissue11,13.

A superficial resemblance between nanomaterials such as CNTs and asbestos has led scientists to challenge the research community to 'Assess whether fibre-shaped nanoparticles present a unique health risk'¹⁴. Published studies have evaluated acute responses to CNTs in cell cultures and the lungs of animal models^{10,15–17}, but the hypothesis that CNTs can behave like asbestos at the mesothelium has not previously been tested. The mesothelial layer is the cell layer that covers the internal surfaces of the pleural (chest) and peritoneal (abdominal) cavities and the exterior surfaces of the organs they contain, lubricating their motion. When cancer occurs in the mesothelium, as is the case

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*Endotoxin detection limit <10 pg ml⁻¹ [†]Metal analysis detection limit <0.1 μg g⁻¹. [‡]The presence of long fibres could not be reliably determined.

	NT _{tang} ;	NT _{tang2}	NT _{long1}	NT _{long2}
Source				
	NanoLab, Inc.	NanoLab, Inc.	Mitsul & Co.	Dr lan Kinloch (University of Manchester)
	tion of morphology (from SEM, TEM an			
	WNTs forming tightly packed	Bundles of intermediate-length	Dispersed bundles and singlets	Regular bundles and ropes or
	l agglomerates, a large proportion of	MWNTs. Often stellate in form with	of long and intermediate-length	MWNTs with a fairly constant
	re in the respirable size range	longer fibres protruding from the	MWNTs, many in the range	length and diameter. Typicali
	with frayed edges of singlet	central tangled agglomerate, a	10-20 µm and longer. Many very short fibres often decorate	single ropes of tubes are mo- than 20 µm in length.
nanotube	FS.	large proportion of which are in respirable size range <5 \mum.	the long fibres.	tilati 20 pari ili lengal.
Diamete	er as supplied by the manufacturer (nr			/
	15±5	15±5	40-50	20-100
Diamete	er as determined by authors (nm, mear	1 + s.e.m.)		
	14.84 <u>+</u> 0.50	10.40 ± 0.32	84.89 <u>+</u> 1.9	165.02 <u>+</u> 4.68
ength:	as supplied by the manufacturer (µm)			
	1-5	5-20	Mean 13	Max 56
Percent	age fibres greater than 15 µm (see Su	pplementary information, Methods, for meti	nodology)	
	‡	‡	24.04	84.26
Percent	age fibres greater than 20 µm (see Su	pplementary information, Methods, for meth	nodology)	
	*	‡	11.54	76.85
Endotox	dn (pg ml ¹)*			
	ND	ND	ND	ND
Soluble	metals (μg g ¹) [†] (see Supplementary in	nformation, Fig. S2, for a full analysis)		
Fe	7.9	13.4	ND [†]	37.3
Cu	5.1	1	1.2	1.2
V	ND†	ND†	0.8	ND
Ni	9.7	5	6.2	6.2
Zn	5.5	7.5	0.7	ND†
Co	3.7	ND†	1.9	3.4

in a proportion of individuals exposed to asbestos, the cancer is termed mesothelioma¹¹. Mesothelioma is almost exclusively found following asbestos exposure and mesothelioma is a particle response unique to fibre-shaped particles¹⁸.

We previously recorded that direct exposure of the mesothelium by intraperitoneal (i.p.) injection of long amosite asbestos fibres resulted in an exaggerated inflammatory response, but there was little or no response to injected short amosite fibres¹⁹. This most likely reflects the extreme sensitivity of the mesothelium to long fibres^{20,21}. We therefore hypothesize that the most important acute response to long CNTs, if they were to behave like asbestos, would be mesothelial injury.

Amosite, or brown asbestos, is an amphibole form of asbestos and the samples used here were prepared for pathology studies undertaken to investigate the role of fibre length²². The long-fibre amosite (LFA) used in this study was South African amosite, which was prepared to have a large proportion of long fibres, and the short-fibre amosite (SFA) sample was prepared from the long-fibre sample by comminution in a ball mill²². In the present study LFA and SFA were used as positive and negative controls, respectively^{19,22}. The four samples of MWNTs were chosen to assess the role of long rigid fibres in stimulating a mesothelial response similar to that caused by LFA. Two of the MWNT samples, NT_{long1} and NT_{long2}, contained a substantial proportion of long straight fibres longer than 20 µm, and the other two samples, NT_{tang1} and NT_{tang2}, consisted of CNTs arranged in low-aspect-ratio tangled aggregates. Full characterization of the MWNT samples and the

derivation of these classifications are provided in Table 1. In addition, a nanoparticulate carbon black (NPCB) sample was used as a non-fibrous graphene control. Transmission electron microscopy (TEM) and optical analysis of NT_{long2} samples confirmed that the nanotube fibres were not all single tubes, but rather were 'wires' or 'ropes' of intertwined single CNTs, rendering them visible by light microscopy in the tissue and cell samples.

Each material was injected in a 50 µg dose into the peritoneal (abdominal) cavity of mice and the cavity systematically washed out at 24 h or 7 days post exposure with physiological saline, in a process termed lavage, to provide a lavageate that contained the cells and molecules present in the peritoneal cavity. The response was assessed by measuring protein levels and cell populations in the lavageate of the peritoneal cavity 24 h later¹⁹. The pathogenic response to long fibres is typified by an alteration in the normal structure of the mesothelial surface to produce a scar-like structure (lesion) called a granuloma. Histological quantification of granulomas on the peritoneal side of the diaphragm²¹ and foreign body giant cells (FBGCs) was carried out after 7 days.

The normal inflammatory response to pathogenic particles or their soluble components is inflammation, with polymorphonuclear leukocyte (PMN) and protein exudation. Only the samples that contained long fibres (LFA, NT_{long1} and NT_{long2}) caused significant PMN (Fig. 1a) or protein (Fig. 1b) exudation. FBGCs (Fig. 1c) and granulomas on the peritoneal side of the diaphragm (Fig. 1d; see also Fig. 2b,c and Supplementary Information, Tables S1 and S2) were also seen with the

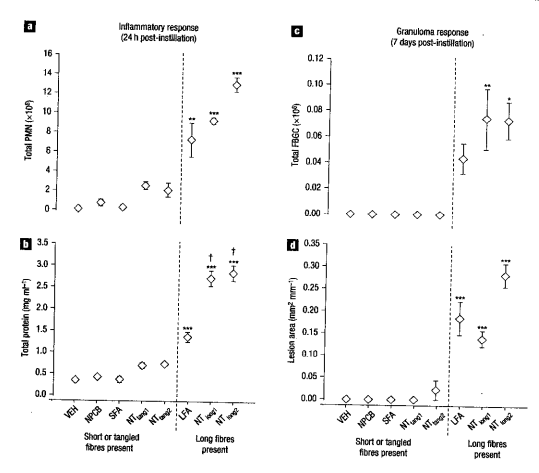


Figure 1 Recruitment of inflammatory cells in the peritoneal cavity after introduction of fibres. Female C57Bl/6 mice were intraperitoneally instilled with 50 μ g of vehicle control (VEH, 0.5% BSA/saline; 0.5 ml), nanoparticulate carbon black (NPCB), short-fibre amosite (SFA), long-fibre amosite (LFA), two curled/tangled MWNT samples of different lengths (NT_{tang1}, NT_{tang2}) and two samples containing long MWNTs (NT_{long1}, NT_{long2}). After 24 h and 7 days post-exposure, the mice were killed and the peritoneal cavity lavaged. a,b, At 24 h, inflammatory response was evaluated using differential cell counts to establish total PMN population (a, mean \pm s.e.m.; VEH, n=3; NPCB, n=3; SFA, n=3; NT_{tang1}, n=3; NT_{tang2}, n=4; LFA, n=6; NT_{long1}, n=4; NT_{long2}, n=3), and total protein (b, mean \pm s.e.m.; VEH, n=5; NPCB, n=4; SFA, n=3; NT_{tang1}, n=6; NT_{tang2}, n=6; LFA, n=6; NT_{long1}, n=8; NT_{long2}, n=6). c, At 7 days, granuloma response was investigated by measuring the total foreign body glant cell (FBGC) population (mean \pm s.e.m.; VEH, n=6; NPCB, n=4; SFA, n=5; NT_{tang1}, n=4; NT_{tang2}, n=4; NT

long-fibre-containing samples. This foreign body response is the normal reaction to indigestible or non-degradable material that macrophages cannot eliminate. The granulomas comprised aggregates of cells containing fibres, most likely macrophages, and also FBGCs, with MWNT/asbestos fibres and associated deposition of collagen within the lesions (Fig. 2b,c). The mesothelial lining on the pleural side of the diaphragm was normal in every case. The PMN inflammation had decreased by 7 days, as the focus of inflammation changed from the peritoneal cavity in general to the granulomas (data not shown). In contrast, particle samples that did not contain detectable long fibres (although they may have contained low-level contamination with long fibres)-NPCB, SFA, NT_{tang1} and NT_{tang2}—failed to cause any significant inflammation at 1 day or giant cell formation at 7 days. A small nonsignificant granuloma response in one of three of the NT_{tang2}-treated mice (see Fig. 1d; see also Supplementary Information, Table S1 and Fig. S3) could have been a consequence of a contamination of long fibres that was so low that they were not detected in the fibre size analysis.

Alternatively, this low-level response could have been caused by some other unidentified component of the NT_{tang2}, or the granulomas could have arisen spontaneously by chance. Although statistical analysis indicated a clear lack of effect from the short MWNT samples, a similar study in a larger group of animals would provide greater confidence in these differences. Neither soluble metals nor endotoxin contamination of long samples were correlated with the greater inflammogenicity and granuloma formation seen with the NT_{long} samples (see Table 1).

To further examine the role of water-soluble components, these were collected from a 50-µg dose of NT_{long1} and NT_{long2} by overnight mixing in 0.5% bovine serum albumin (BSA)/sterile saline vehicle. The soluble components produced no significant inflammatory effects 24 h after injection into the mouse peritoneal cavity (see Supplementary Information, Fig. S2). Neither can levels of total metals explain the differences between short and long responses seen with nanotube samples (see Supplementary Information, Fig. S1). Transition metals have



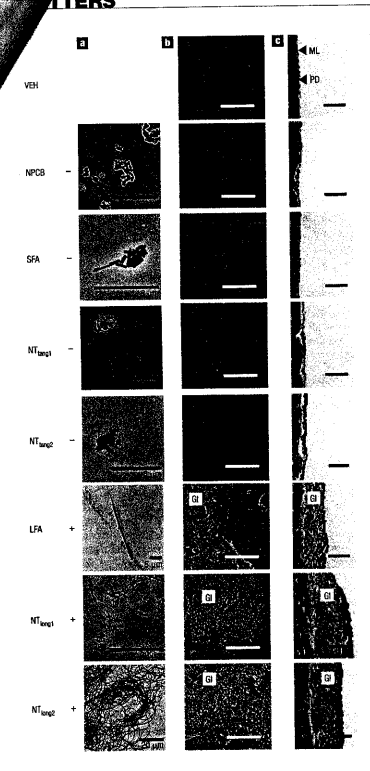


Figure 2 Effect of the particle/fibre on diaphragms after 7 days. a, TEM images show the presence (-1) or absence (-) of long fibres in the samples used. Female C57BI/6 mice were injected i.p. with 50 μg of sample, killed after 7 days, and the diaphragms excised and prepared for visualization. b,c, SEM images (b) and haematoxylin and eosin histology sections (n=3) (c) of the diaphragms show the presence of granulamatous inflammation (GI) in mice exposed to LFA, NT_{long1} and NT_{long2}. A small granuloma response in one of the three mice treated with NT_{lang2} (see Supplementary Information, Tables S1 and S2, for raw data and statistical analysis) was observed. The muscular portion of the peritoneal diaphragm (PD) and the mesothelial layer (ML) are aligned to show granulomatous inflammation at the peritoneal aspect of the diaphragm surface. Scale bars in b: 200 μm . Scale bars in c: 50 μm .

been implicated in CNT-mediated stress in an earlier published study on nanotubes in vitro²³, but neither soluble nor total metals can explain the differences in peritoneal response seen here with long and short MWNTs. There is a remote possibility that some unmeasured metal or other component could contribute to this difference but, in the opinion of the authors, it is unlikely. Although our data show that short CNTs do not mimic the behaviour of long asbestos, our data cannot preclude the possibility that short CNTs are harmful by conformation to some other paradigm that was not addressed here, such as intrinsic toxicity as particles, as opposed to fibres.

In attempting to phagocytose or engulf a fibre longer than the length they can completely enclose, the specialized 'engulfing cells', macrophages, are chronically stimulated to release mediators that cause inflammation^{24,25}. In this inflammatory milieu the conditions, including the cytokine environment, that encourage macrophage fusion are produced, leading to giant cell formation (Fig. 3c,f). The macrophages that are undergoing frustrated phagocytosis are not necessarily the ones that form giant cells, as shown by the fact that macrophages are stimulated to undergo formation into giant cells in vitro with a combination of cytokines alone²⁶. So, although we have no quantitative data on the role of frustrated phagocytosis, images from the lavage of animals exposed to LFA and long MWNTs (Fig. 3a,d) show that frustrated phagocytosis did occur. No FBGCs were found in the lavage of animals exposed to NPCB, SFA, nor the short, tangled MWNT samples, and complete phagocytosis was evident in macrophages from the lavage (see Fig. 3b,e for SFA and NT_{tang1}, respectively).

Our aim was to investigate commercial samples of CNTs to obtain real-world relevance and compare pathogenicity and we showed clear differences between long and short/tangled MWNTs. However, there was an attendant problem in that there are differences in the source, preparation and purification of different commercial CNTs and therefore potential differences in physicochemistry and contaminating metals. For example, the two short tangled samples were acid-treated and this may have resulted in alterations. We have addressed this here and largely discounted factors other than length, but another strategy to critically address the role of length is to mill long CNTs and compare the unmilled (long) with the milled (short) fibres, so minimizing differences except for length. Short CNTs may be pathogenic by virtue of being particles, and that would not have been detected in the assays used here, which are sensitive only to asbestos/fibre-type effects.

The discrimination in terms of asbestos-like hazard between short/tangled and long MWNTs shown after low-dose injection of fibres in our study emphasizes the importance of choosing the correct assay. In this case the assay was based on the uniqueness of the mesothelial response to long fibres. We observed that long MWNTs produced inflammation, FBGCs and granulomas that were qualitatively and quantitatively similar to the foreign body inflammatory response caused by long asbestos. In contrast, NPCB did not cause peritoneal inflammation, highlighting the fact that fibrous shape dominates over simple graphene chemistry in effects on the mesothelium.

The fact that CNTs can be produced in a morphology that implies an asbestos-like hazard, although unwelcome, was predicted on the basis of the fibre paradigm^{4,12}. The single published study on CNTs in workplace air²⁷ found that airborne particles that resemble the tangled MWNT samples shown here have low potency. We note that two of the three commercially obtained samples of MWNTs used here (NT_{tangl} and NT_{tang2}) had similar short/tangled morphologies due to their curled/tangled structure. This morphology showed little potency in the assays and so these would not be predicted to behave like asbestos.

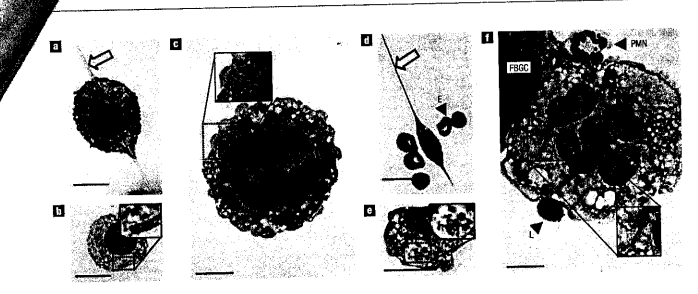


Figure 3 Effect of fibre length on phagocytosis by peritoneal macrophages. a,b, Histological sections show incorporation of LFA (a, arrow) leading to 'frustrated phagocytosis', but SFA (b, see inset) is successfully phagocytosed. c, Representative image of an FBGC after injection of LFA containing short fragments of fibre (see inset). d, Like LFA, NT_{long2} also leads to frustrated phagocytosis (E-erythrocytes). e, in contrast, NT_{tang1} can be readily phagocytosed (see inset). f, FBGC is also present after injection of NT_{long2} (see inset for internalized fibres) (PMN, polymorphonuclear leukocyte; L, lymphocyte). All images are shown at $\times 1,000$ magnification with a 5 μ m scale bar.

Our data demonstrate that asbestos-like pathogenic behaviour associated with CNTs conforms to a structure-activity relationship based on length, to which asbestos and other pathogenic fibres conform. Exposure of the mesothelium, however, is based upon the caveat that all materials tested shared a high level of biopersistence, allowing sufficient time for migration through the lung to the mesothelium. As such, our study does not address whether CNTs would be able to reach the mesothelium in sufficient numbers to cause mesothelioma following inhalation exposure. Although the study suggests a potential link between inhalation exposure to long CNTs and mesothelioma, it remains unknown whether there will be sufficient exposure to such particles in the workplace or the environment to reach a threshold dose in the mesothelium. Intraperitoneal injection of long asbestos into the peritoneal cavity of rodents has been demonstrated to cause mesothelioma in the long term²². However, our study did not address whether the mice exposed to long CNTs that developed inflammatory and granulomatous changes would go on to develop mesotheliomas. Neither did our study rule out the possibility that short CNTs may have some intrinsic pathogenicity by virtue of their particulate nature, which would not have been detected in the assays used here, which are specific for fibre effects. These are research questions that must be addressed with some urgency before the commercial use of long CNTs becomes widespread.

METHODS

A panel of four different samples of MWNTs and three different control particles (LFA, SFA and NPCB) was used. The MWNT panel comprised three commercially available nanotubes and one sample produced in an academic research laboratory. The nanotube samples used were a sample of long MWNTs (NT_{long1}; Mitsui & Co.), a second MWNT sample of largely long fibres (NT_{long2}; University of Manchester), and two curled/tangled NT samples either cut (using a proprietary process) to form predominantly short fibres (NT_{tang1}) or with their original length (NT_{tang2}) (NanoLab). The NPCB used was Printex 90 (Degussa). The classification of samples into NT_{long} or NT_{tang} was established from the physical characteristics observed using scanning electron microscopy (SEM), TEM and light microscopy (only TEM images are shown), where appropriate,

to fully describe each sample (Table 1). The diameter of both individual fibres and (in the case of NT $_{\rm tang1}$ and NT $_{\rm tang2}$) agglomerate size as well as a length size distribution analysis of the fibres and particles were established. In keeping with World Health Organization (WHO) guidelines we counted as a fibre only those particles with an aspect ratio greater than 3:1 and a length $> 5~\mu m$.

Inductively coupled plasma mass spectrometry (ICP-MS) analysis was used to quantify contaminating metals (Table 1; see Supplementary Information, Fig. S1 for a full metals analysis), and the presence of bacterial contamination of the particle panel was established by measuring endotoxin levels (Table 1). Samples were prepared for in vivo use by ultrasonication in a sterile 0.5% BSA/saline solution and were administered by injection into the peritoneal cavity of female C57Bl/6 mice (aged 8 weeks) at a dose of 100 µg ml⁻¹ (0.5 ml; total dose 50 µg per mouse). After 24 h and 7 days the mice were killed by cervical dislocation and the peritoneal cavity lavaged using three 2 ml washes of ice-cold sterile saline, which were pooled together on ice. The 24-h timepoint was chosen based on our experience from previous studies, which indicated that at this timepoint we would be able to discriminate between different particles in their ability to cause inflammation. The 7-day timepoint was chosen to allow time for fibrosis to develop in the granulomas at the diaphragm. The lavage fluid was centrifuged to separate the cellular fraction, and the supernatant protein content was established using a BCA protein assay (VEH, n = 5; NPCB, n = 4; SFA, n = 3; NT_{tang1}, n = 6; NT_{tang2}, n = 6; LFA, n = 6; NT_{long1}, n = 8; NT_{long2}, n = 6). Cyto-centrifugation preparations were made using the isolated cells, and differential cell counts were performed to establish leukocyte populations (VEH, n = 3; NPCB, n = 3; SFA, n = 3; NT_{tang1}, n = 3; NT_{tang2}, n = 4; LFA, n = 6; NT_{long1} , n = 4; NT_{long2} n = 3). At 7 days the diaphragm was carefully removed from the mice and fixed, sectioned, and haematoxylin and eosin stained for gross pathology (VEH, n = 4; NPCB, n = 3; SFA, n = 3; NT_{tang1}, n = 3; NT_{tang2}, n=3; LFA, n=3; NT_{long1}, n=4; NT_{long2}, n=3). The granulomatous lesions present with each section was quantified using serial images taken at ×100 magnification using QCapture Pro suftware (Media Cyberbernetics) and the mean lesion area and length of the complete diaphragms (see Supplementary Information, Fig. S3) established using Image-Pro Plus software (Media Cybernetics). Diaphragms removed for surface analysis by SEM were fixed, osmified and gold-sputter-coated for viewing under SEM. All data are expressed as a mean ± s.e.m. and analysed using one-way analysis of variance (ANOVA). Multiple comparisons were analysed using the Tukey-HSD method and in all cases, values of P < 0.05 were considered significant. (See Supplementary Information for a full supplementary methodology.)

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Author contributions

C.A.P., R.D. and K.D. initiated, designed, directed and performed all experiments and took responsibility for planning and writing the manuscript. LK. manufactured and provided the MWNT sample, NT_{long2}, and provided specialist input regarding CNT production and morphology, W.A.H.W. aided in the interpretation of the histological sections and contributed to the writing of the manuscript. A.M., A.S., S.B., V.S. and W.MacN. provided intellectual input and contributed to the writing of the manuscript.

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