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Cancer Research

2 Q1 S100A4 Elevation Empowers Expression of 3 Q2 Metastasis Effector Molecules in Human Breast 4 Cancer R

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7 Abstract

Many human glandular cancers metastasize along nerve tracts, 8 9 but the mechanisms involved are generally poorly understood. 10The calcium-binding protein S100A4 is expressed at elevated 11 levels in human cancers where it has been linked to increased 12invasion and metastasis. Here we report genetic studies in a 13Drosophila model to define S100A4 effector functions that mediate 14metastatic dissemination of mutant Ras-induced tumors in 15 the developing nervous system. In flies overexpressing mutant Ras^{Val12} and S100A4, there was a significant increase in activation 1617of the stress kinase JNK and production of the matrix metallo-18proteinase MMP1. Genetic or chemical blockades of JNK and 19 MMP1 suppressed metastatic dissemination associated with -33

34 Introduction

35 Certain tumor cells have a propensity to invade the neighboring 36 tissue and eventually establish new secondary tumors or metas-37 tases while others cannot (1, 2). These results suggest that a 38 specific set of genes, different from those involved in the produc-39 tion of the neoplasia, are involved in promoting a complex series 40 of steps to form metastases (3). The protein products of such 41 genes have been termed metastasis-inducing proteins (MIP). One 42such gene/protein is S100A4 (4), a member of the S100-calcium-43binding protein family (5). Although S100A4 cannot promote 44 tumor formation directly, it can stimulate the remaining steps in 45the metastatic cascade in model rodent systems by combining with oncogenes such as Ras^{Val12} and neu (4, 6). Moreover, S100A4 4647is overexpressed in human primary tumors and is associated with 48the premature death of patients with different types of metastatic 49carcinomas, including those from the breast (7), oral mucosa, 50bladder, pancreas, prostate, colorectum, esophagus, lung, stom-51ach, and thyroid glands (8). The elevation of S100A4 can trigger 52multiple biological functions, including cell migration, invasion, 53extracellular matrix remodeling, and angiogenesis (8, 9). How-

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S100A4 elevation, defining required signaling pathway(s) for 21S100A4 in this setting. In clinical specimens of human breast 22cancer, elevated levels of the mammalian paralogs MMP2, MMP9, 2324and MMP13 are associated with a 4- to 9-fold relative decrease in 25patient survival. In individual tumors, levels of MMP2 and MMP13 correlated more closely with levels of S100A4, whereas 2627MMP9 levels correlated more closely with the S100 family 28member S100P. Overall, our results suggest the existence of evolutionarily conserved pathways used by S100A4 to promote 29metastatic dissemination, with potential prognostic and thera-30 31peutic implications for metastasis by cancers that preferentially exploit nerve tract migration routes. Cancer Res; 1-12. ©2016 AACR. 32

ever, it is unknown what are the biologically relevant molecular events from the plethora triggered by S100A4 in cultured mammalian cells (10). To generate a genetically tractable experimental model to investigate the molecular events triggered by S100A4, we have for the first time expressed human S100A4 in the fruit fly, Drosophila melanogaster by targeting its expression to the developing eye lobes (11) and not elsewhere in the brain or CNS (12, 13). Drosophila has conserved signal transduction pathways for cell cycle, growth control, and cell-to-cell communication (14) and a larval phase of only a few days, which can be interrogated by inhibitory chemicals applied directly to the growth medium. In addition, 70% of human cancer genes are conserved in the Drosophila model, but importantly none of the S100 family proteins are present (15). Overexpression of oncogenic Ras (*Ras-Val12*) causes the formation of tumors in the epithelial tissues of Drosophila (16), and these can be transformed into a malignant phenotype by disruption of suppressor genes such as scribble

(scrib) and lethal2 (17). In this new model we have for the first time generated transgenic flies capable of conditionally-expressing the open-reading frame of humanS100A4 under GAL4/UAS-control (18). We show, after multiple crosses, that the S100A4 gene is required to disseminate Ras^{Val12} tumor cells from the optic lobes to the ventral nerve cord (VNC) and further afield in fly larvae. The combination of Ras^{Val12} and loss of *scrib* in *Drosophila* activates the JNK pathway and this activation induces the matrix metalloproteinase MMP1 to allow dissemination of the cancer cells in the Ras oncogenic system (16). Therefore, we have assessed whether c-Jun and *Drosophila* MMP1 are downstream targets for promoting dissemination in our $Ras^{Val12}/S100A4$ novel larval model using fly genetics and inhibitory chemicals, and whether there is a uniquely similar association between S100A4 and mammalian MMPs in human breast cancer.

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Note: Supplementary data for this article are available at Cancer Research Online (http://cancerres.aacrjournals.org/).

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89 Materials and Methods

90 Expression of S100A4 in Drosophila melanogaster

91HumanS100A4 wild-type (S100A4wt; ref. 19) and inactive mutantS100A4 Δ 2 (20) were cloned and expressed in transgenic 9293 flies as described in Supplementary Methods. Stable transgenic 94 lines were checked for S100A4 expression by crossing with da-95GAL4 flies (18) and Western blotted. Resultant S100A4wt and 96 S100A4 Δ 2 progeny produced (mean \pm SE) similar 7.7 \pm 0.6 ng 97 and 8.8 \pm 0.7 ng S100A4 protein per 20 flies, respectively, 98 compared with undetectable levels (<0.1 ng) in parental con-99 trols [Student t test (STT); P = 0.29]. Remaining details are in 100 Supplementary Methods. All initial Drosophila strains were 101 described previously (21), remaining details are in Supplemen-102tary Methods. The flies were maintained in standard yeast agar 103medium at 25°C in a 12-hour light-dark incubator (21).

104 Metastatic assay

105The eyeless-FLP-induced recombination of the FRT-flanked y 106linker in Act>y>GAL4 results in reconstitution of Actin-GAL4 and 107 expression of UAS-GFP, and other UAS elements, in the devel-108 oping eye (22). Dissemination of GFP from its original site of eye-109 antennal discs to VNCs was scored for each genotype/experimen-110tal condition on a scale of 0 to 3 (16). GFP localized in the optic 111 lobes scored zero (stage 0), GFP on one side of the VNC scored 1 (stage I), on two sides of VNC scored 2 (stage II), and dissemi-112 nation further into the VNC scored 3 (stage III). Average stage 113 114score of metastasis (ASSM) ±SE was recorded for each genotype/ 115experimental condition. Fluorescent staining is described in Sup-116 plementary Methods. Confocal images of GFP were captured (21) 117and analyzed using ImageJ software (23), as described in Sup-118plementary Methods. Corrected integrated fluorescence intensity 119 (CIFI) = integrated intensity - (area of selected background brain \times mean fluorescence density of background) (23). Mean 120 121CIFI \pm SE was recorded.

122 Western blot analysis

This is described in Supplementary Methods. To correct for any 123124loading differences, original intensity of each band was divided by 125that of actin. Intensity of each band for a particular larval group was then expressed as a ratio of that in the Ras^{Val12} larvae. Mean 126value of three experiments $\pm SE$ was recorded. To ensure the 127128intensity of band signals lay within the linear range, a plot of band intensity against µg GAPDH was drawn ($y = 43947 \times$ 129130-42398, $r^2 = 0.997$) and band intensity for any protein outside the linear range was excluded from the data and if necessary the gel 131 132was rerun with higher or lower levels of total protein.

133 Drug treatment

134Inhibitors, JNK-IN-8 (kindly provided by Nathanael S. Gray, 135Harvard Medical School, Boston, MA; ref. 24) and batimastat (cat. 136no.: SML0041; Sigma-Aldrich; ref. 25) were added directly from 1371 mg/mL stock dissolved in DMSO to larval medium preheated to 13855-60°C. Same concentration of DMSO was added to controls 139without inhibitors. The drugs were incubated with the larvae 140 continuously until harvesting at the third instar stage, equivalent to 7 days. 141

142 Statistical analyses

143 The significance of the difference between two categorical 144 groups for each genotype, those with and those without

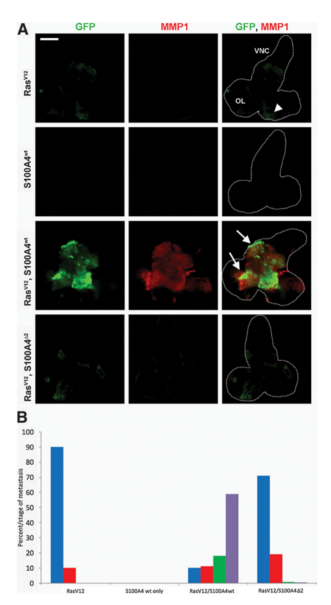


Figure 1.

A, Fluorescent images of GFP and MMP1 in larval CNS of different male recombinant Drosophila. CNS was dissected from third instar larvae of Drosophila with the following backgrounds: Ras^{Vall2}(RasV12) only; S100A4 wild type (S100A4wt) only; Ras^{Val12}, S100A4 wild type (RasV12/S100A4wt); and Ras^{Val12}, S100A4 Δ 2 (RasV12/S100A4 Δ 2). Representative CNS images show green fluorescence due to endogenous GFP, red fluorescence due to fluorescently labeled antibodies to matrix metalloproteinase1(MMP1), and merged fluorescent images are due to GFP and anti-MMP1. The outline of the relevant structures of the brain including optic lobes (OL) and ventral nerve cord (VNC) are indicated by the broken white line. The region which clearly expresses MMP in Ras^{Val12}, S100A4 transgenics (arrows), and the same region in Ras^{Val12} transgenics (arrow head) are shown (Scale bar, 100 µm). B, Histogram of resultant recombinant larvae. The percentage (percent) larvae with different stages (0-III) of metastasis from the optic lobes to the ventral nerve cords (VNC) is shown for the recombinant Drosophila. The VNC of at least 50, third instar larvae were scored for the extent of metastasis on a sliding scale (Materials and Methods): from stage 0 (blue), stage I (red), stage II (green), and stage III (purple). Larvae containing Ras^{Val12}/S100A4 were significantly different from these containing $Ras^{Va/12}$ alone, S100A4 alone, and $Ras^{Va/12}/S100A4\Delta 2$ (Fisher exact test P < 0.0001) and between larvae containing Ras^{Vall2} and $Ras^{Vall2}/$ $S100A4\Delta 2 (P = 0.012)$

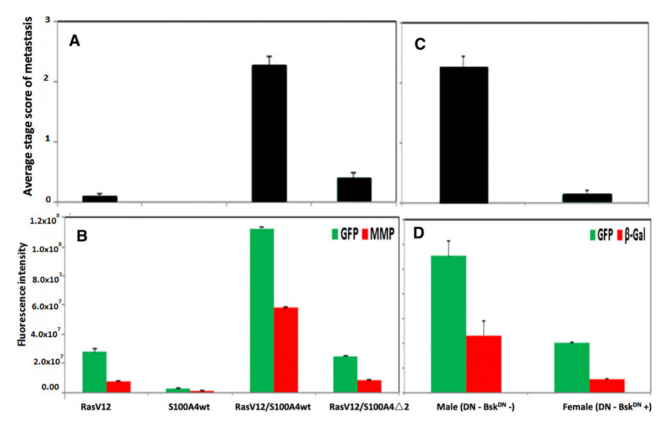


Figure 2.

The provides a set of the primary tumor in the primary tumor in the primary tumor in the optic lobe spreading to the VNC is shown for male recombinant *Drosophila* with genetic backgrounds of: *Ras^{Val12}/S100A4* wild *type* only (S100A4wt), *Ras^{Val12} plus S100A4 wild type* (RasV12/S100A4wt), and *Ras^{Val12} plus S100A4mutant* (RasV12/S100A4Δ2). At least 50 larvae were scored (Materials and Methods) and results are expressed as mean ± SE. Both double transgenic flies were significantly different from flies with *Ras^{Val12}/S100A4wt* from *Ras^{Val12}/S100A4* genotype (P = 0.0001). **B**, Fluorescence intensity of CNS images of *Ras^{Val12}/S100A4* flies. Endogenous fluorescence from GFP (green) and from exogenously-added labeled antibody to MMP1 (red) were recorded. CIFI of images of the dissected CNS from the same larvae in A were computed as described in "Methods." Mean ± SE is shown. For GFP green fluorescence, *Ras^{Val12}/S100A4wt* versus *Ras^{Val12}/S100A4* (STT.P < 0.0001); *Ras^{Val12}/S100A4* (P = 0.49). For MMP-1 red fluorescence, *Ras^{Val12}/S100A4wt* versus *Ras^{Val12}/S100A4* (STT.P < 0.0001); *Bas^{Val12}/S100A4* (P = 0.49). For MMP-1 red fluorescence, *Ras^{Val12}/S100A4wt* versus *Ras^{Val12}/S100A4* (STT.P < 0.0001); *Bas^{Val12}/S100A4* (P = 0.49). For MMP-1 red fluorescence, *Ras^{Val12}/S100A4wt* versus *Ras^{Val12}/S100A4* (STT.P < 0.0001); *Bas^{Val12}/S100A4* (P = 0.49). For MMP-1 red fluorescence, *Ras^{Val12}/S100A4wt* versus *Ras^{Val12}/S100A4wt* only, on *Ras^{Val12}/S100A4A2* (STT.P < 0.0001); but Ras^{Val12}/S100A4A2 (P = 0.49). For MMP-1 red fluorescence, *Ras^{Val12}/S100A4wt* versus *Ras^{Val12}/S100A4Wt* versus *Ras^{Val12}/S100A4Wt* only, or *Ras^{Val12}/S100A4A2* (STT.P < 0.0001); but Ras^{Val12}/S100A4A2 (P = 0.5). **C**, Average stage of metastatic spread in male and female JNK-suppressed *Ras^{Val12}/S100A4Wt* in which the *Bsk^{DN}* dominant suppressor of JNK is expressed only in female flies. At least 20 larvae were scored as in **A**. Results are expressed as mean ± SE and there was a highly s

147metastases was determined by Fisher exact test, recording two-148sided values of P. The significance of the difference in ASSM, in149CIFI for GFP and MMP1, and in mean corrected intensity of each

- 150 protein band in Western blots were calculated using two-sided
- 151 Student *t* test (Stats Direct). Differences considered significant 152 when P < 0.05.

153 Patients and specimens

154A retrospective study was undertaken using samples of 183155primary tumors from unselected breast cancer patients, as156described previously (26, 27). Ethical approval was obtained157from NRES Committee, North West REC.Ref. 12/NW/0778, Pro-158tocol no. UoL000889, IRAS no. 107845. Samples were preserved

in neutral buffered-formalin and embedded in paraffin-wax, as 160 described previously (7). 161

IHC staining

This is described in Supplementary Methods. Western blots of 163breast cell lines verified the specificity of all three mAbs to MMPs 164yielding apparent molecular weights of 73,99,75 kDa for secreted 165latent MMP2, 9, 13, respectively, consistent with those reported 166recently (28). Remainders were verified previously (27). IHC-167stained sections were analyzed and scored (7, 26, 28, 29), as 168recorded in Supplementary Methods. Association of staining for 169MMP2, 9, and 13 with patient survival time is reported in 170Supplementary Methods. 171

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174 **Results**

Cooperation of Ras^{Val12} and S100A4 in producing metastases in recombinant *Drosophila*

177The brain containing the CNS was dissected from at least fifty, 178third instar larvae of different recombinant Drosophila. Male larvae with the genetic background Ras^{Val12} alone (Ras^{Val12} larvae) 179180 produced GFP-fluorescent tumors almost exclusively in the eye 181 lobes (Fig. 1A) of 48 of 53 cases, with only 5 of 53 cases extending 182into one or the other side of the VNC (Fig. 1B). Extent of metastasis 183was semiquantified as described in Materials and Methods to produce an ASSM. There were no GFP-tumor deposits in the eye 184 185lobes or elsewhere in S100A4 larvae (Figs. 1A and B and 2A). The Ras^{Val12}/S100A4wtrecombinant larvae produced metastasis to the 186 187VNC (Fig. 1C) in a significantly higher number of 53 of 59 cases [Fisher exact test, P (FET.P) < 0.0001; Fig. 1B], increasing the ASSM 188 by a significant 24-fold over Ras^{Val12}larvae [Student t test P (STT.P) 189190< 0.0001; Fig. 2A]. There was also extensive metastasis to other 191organs, particularly to the gut and gonads (Supplementary Fig. S1). The Ras^{Val12}/S100A4 Δ 2 inactive mutant larvae (Materials and 192Methods) produced a significantly lower number of 16 of 56 with 193metastasis to the VNC (FET.P < 0.0001; Fig. 1B), with significant 1945.7-fold reduction in ASSM compared to Ras^{Val12} larvae (STT.P = 1950.0001; Figs. 1A and 2A). The CIFI of GFP (Materials and Meth-196 ods) for images taken of the dissected CNS of Ras^{Val12}larvae 197was increased by a significant 4.1-fold in Ras^{Val12}/S100A4wt 198larvae (STT.P < 0.0001; Fig. 2B), but there was no significant 199difference in CIFI of Ras^{Val12}/S100A4A2mutant larvae compared 200 with Ras^{Val12} larvae (STT.P = 0.49; Fig. 2B). 201

202Quantification of Ras, GFP, and S100A4 levels by Western blot203analysis

204Antibodies to S100A4 detected a specific band of the correct apparent molecular weight of 9 kDa in all fly lines containing the 205206S100A4wt or S100A4 Δ 2 mutant gene, but no corresponding band in larvae containing Ras^{Val12} alone (Supplementary Fig. S2). 207Larvae containing the $Ras^{Val12}/S100A4wt$ and $Ras^{Val12}/S100A4\Delta 2$ 208genes produced a significant increase in Ras and a similar increase 209210in GFP over that in larvae containing Ras^{Val12} alone (STT.P < 2110.001; Table 1). Protein bands of Ras and GFP were observed at 212the correct molecular weights (21 and 27 kDa, respectively; Supplementary Fig. S2 and Table 1). There was also highly 213

Q6 Table 1. Quantification of Western blots of different Drosophila lines

significant increases of 220 \pm 5- and 85 \pm 11-fold in S100A4 protein in larvae containing the *Ras^{Val12}/S100A4wt* and *Ras^{Val12}/S100A4* Δ 2 genes, respectively (*P* < 0.0001; Table 1), when normalized to GFP. Thus, there is a significant association of expression of active S100A4 and metastasis in this model system.

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Increased levels of activated JNK and MMP1 in Ras and S100A4-overexpressing larvae

Levels of JNK in Ras^{Val12} and Ras^{Val12}/S100A4wt larvae were not significantly different in Western blots analysis (STT.P = 0.50; Table 1). However, levels of activated phospho-JNK and MMP1 at the reported molecular weights of 46 and 52 kDa, respectively (30), rose significantly by 13.1 \pm 0.6- and 3.8 \pm 0.1-fold, respectively, when normalized to GFP, in RasVal12/ S100A4wt compared to Ras^{Val12} larvae (P < 0.0001; Supplementary Fig. S2 and Table 1). There was no significant increase in phospho-JNK, JNK, and MMP1 in Ras^{Val12}/S100A4 Δ 2 compared to Ras^{Val12}larvae. In S100A4wt larvae alone, the levels of phospho-INK, INK, and MMP1 were significantly lower (P < 0.0001, P < 0.0001, P =0.02; Supplementary Fig. S2 and Table 1), probably reflecting the absence of any primary tumor (Figs. 1A and B and 2A). There was also a modicum of red fluorescence for MMP1 in the eve lobes of Ras^{Val12}larvae (Fig. 1A and B), which rose significantly in Ras^{Val12}/ S100A4wt larvae (P < 0.0001; Fig. 2B) showing extensive staining of the VNC (Fig. 1A and B). There was no significant difference in CIFI for Ras^{Val12} and $Ras^{Val12}/S100A4\Delta 2$ larvae (Fig. 2B).

Activated JNK and MMP are downstream effectors in Ras and S100A4-overexpressing larvae

To determine the requirement for JNK signaling in the meta-242static phenotypes, we expressed dominant-negative JNK encoded by *basket* (Bsk^{DN}), together with Ras^{Val12} and S100A4. When 243244female and male siblings with and without Bsk^{DN}, respectively 245(Materials and Methods) were examined, 15/15 male, but only 2/ 24615 female larvae produced extensive metastases to the VNC (FET.P 247< 0.0001; Supplementary Fig. S3). ASSM and CIFI were reduced by 248a significant 17- and 2.8-fold in female larvae, respectively (STT.P 249< 0.0001; Fig. 2C and D). The expression of a genetically-engi-250neered marker of JNK activity, puc-LacZ was followed by its 251induction of β-galactosidase (Materials and Methods; Supple-252mentary Fig. S3). The CIFI for red fluorescent antibody to β-galac-253tosidase fell significantly by 4.7-fold in females (P = 0.004; 254

	Mean relative abundance ^b				
Antibody to ^a	Ras ^{Val12}	S100A4wt	Ras ^{Val12} /S100A4wt	Ras ^{Val12} /S100A4∆2	
(A) Ras	1 ± 0.05	0.0097 ± 0.001	$3.56\pm0.08^{\circ}$	$1.96 \pm 0.16^{\circ}$	
(B) GFP	1 ± 0.04	0.011 ± 0.002	3.50 ± 0.05^{c}	$1.60 \pm 0.10^{\circ}$	
(C) S100A4	1 ± 0.06	9.62 ± 2.1	742 ± 19^{d}	136 ± 18^{d}	
(D) P-JNK	1 ± 0.01	$0.19\pm0.02^{\text{e}}$	40.5 ± 0.2^{d}	1.32 ± 0.08	
(E) Total JNK	1 ± 0.04	$0.23\pm0.01^{\text{e}}$	1.15 ± 0.07	1.04 ± 0.10	
(F) MMP	1 ± 0.04	$0.40\pm0.07^{\rm f}$	13.1 ± 0.3	0.498 ± 0.001	
MMP	1 ± 0.04	nd	$2.17\pm0.08^{\rm g}$	nd	

Abbreviation: nd, not determined.

^aTen µg protein larval extracts were treated with the antibody shown in Western blots of Supplementary Fig. S2.

^bMean relative abundance after scanning the blots by densitometry (Materials and Methods) and the area under the peak corresponding to each protein was first normalized to that of actin and then ratioed to the level of that protein in the Ras^{Vall2} male larvae which was arbitrarily set at 1. Mean relative abundance \pm SE from three separate experiments.

^cStudent *t* test P < 0.001 over $Ras^{Va/12}$ male larvae.

^dStudent t test P < 0.0001 over Ras^{Val12} male larvae or S100A4wt male larvae.

^eStudent *t* test P < 0.0001 over Ras^{Vall2} male larvae.

fStudent t test P = 0.02 over Ras^{Vall2} male larvae.

⁹Student *t* test P < 0.0001, for female over male larvae.

257 Supplementary Fig. S3 and Fig. 2D). In Western blots analysis, the 258 level of MMP1 protein normalized to that in male Ras^{Val12}larvae 259 fell 6.0-fold from 13.1 ± 0.3 to 2.17 ± 0.08 in male versus female 260 larvae (P < 0.0001; Table 1).

261When increasing concentrations of the JNK-IN-8 inhibitor (24) were added to male Ras^{Val12}/S100A4wt larvae, there was a signif-262263icant fall in ASSM of 2.5-fold for 1 µmol/L (STT.P < 0.0001), but thereafter a more gradual stepwise decline; the overall fall being 2642655.8-fold (*P* < 0.0001; Supplementary Fig. S4A and S4B; Fig. 3A). 266 There was a similar significant decline in CIFI for GFP of 2.7-fold 267(P = 0.002) for 1 µmol/L inhibitor and then successive significant 268decreases; the overall fall being 11.4-fold (P < 0.0001) (Fig. 3A). 269There was also a similar significant decline in CIFI for antibodies 270to endogenous MMP1 upon addition of 1 µmol/L JNK-IN-8 271(Supplementary Fig. S4A and S4B; P = 0.0005), then further 272successive significant decreases; the overall fall being 8.3-fold 27:Q7 (*P* < 0.0001; Fig. 3A).

274When increasing concentrations of the inhibitor of MMP activity, Batimastat (25) was added to Ras^{Val12}/S100A4wt larvae, there 275were significant falls in ASSM of 3.4-fold for 5 μ mol/L (P < 2762770.0001), but thereafter the decline was more gradual; the overall fall being 16.4-fold (P < 0.0001; Supplementary Fig. S4C and 278279S4D; Fig. 3B). There was a similar significant decline in CIFI for 280 GFP of 3.5-fold (P = 0.0002) for 5 µmol/L inhibitor and then 281successive nonsignificant decreases. The overall fall was 8.4-fold 282 (P < 0.0001; Fig. 3B). There was also a rapid significant decline 283in CIFI for antibodies to endogenous MMP1 upon addition of 2845 µmol/L batimastat (Supplementary Fig. S4C and S4D) of 5.2-285fold (P = 0.028), then nonsignificant successive falls; the overall 286fall being 10.3-fold (*P* < 0.0001; Fig. 3B). Thus, a definite pathway 287has been established between S100A4 and MMP1 for induction of 288metastasis in this model system.

Association of MMPs with patient survival time in human breastcancer

Next, we investigated the relationship in human breast cancer 291292between the more commonly-occurring, mammalian MMPs, 293MMP2, 9, 13, and patient demise as a result of metastatic cancer 294(31). On examination of 183 breast carcinomas for IHC for these 295three MMPs, 32% to 67% contained carcinoma cells which were 296negatively stained (<1% carcinoma cells stained), 19% to 26% 297were borderline stained (1-5% carcinoma cells stained), and the 298rest (15-47%) were stained to varying degrees (Fig. 4 and Sup-299plementary Fig. S5; Supplementary Table S1). There were also 300some reactive stromal cells, mainly myofibroblasts, macrophages, 301 and neutrophils which stained (Fig. 4). Assessment of staining 302 class was made only for the malignant cells. Staining for individual MMPs was abolished by prior incubation of each antibody 303 304with the requisite MMP (Supplementary Fig. S5).

305 To determine whether there was any association between staining for the separate MMPs and of survival of patients, 306 307 Kaplan-Meier survival curves were plotted for different staining 308 groups. Overall, there was a significant difference in staining for 309 each MMP (Wilcoxon Gehan Statistics, P < 0.001). However, the largest significant differences occurred between the (\pm) and (+)310311staining groups for MMP2, 9, and 13, respectively (Supplemen-312tary Table S2). The 183 patients were therefore separated into two 313 categorical groups using a cutoff of 5% stained carcinoma cells for 314 each MMP. Only 11 \pm 4% survived with positively stained 315tumors, compared to $81 \pm 4\%$ with negatively stained tumors 316 for MMP2; 10 \pm 6% vs. 58 \pm 4% for MMP9; and 22 \pm 5% versus

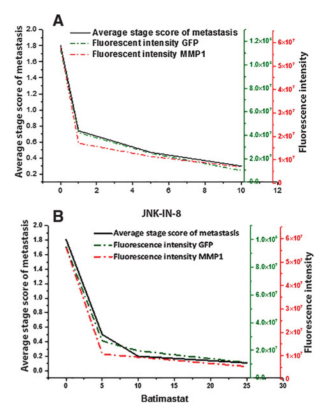


Figure 3.

Tumor dissemination in recombinant flies treated with either JNK-IN-8 (A) or Batimastat (B). Drosophila larvae with genetic background of Ras^{Val12}/ S100A4wt were fed either (A) 0, 1, 5, or 10 µmol/L of the JNK inhibitor JNK-IN-8 or (**B**) 0, 5, 10, or 25 μ mol/L of the MMP1 inhibitor Batimastat in their medium (Materials and Methods). At least 20 larvae were scored and ASSM was computed as described in Materials and Methods. These same larvae were dissected, stained, and scored for endogenous green fluorescence from GFP and for red fluorescence from exogenously-added labeled antibody to MMP1. The CIFI was computed as described in Materials and Methods. Results are shown as mean \pm SE. For ASSM, transgenic larvae fed 0 µmol/L of inhibitor were significantly higher than for larvae fed 1. 5. and 10 umol/L JNK-IN-8 or for larvae fed 5, 10, and 25 $\mu mol/L$ batimastat (STT.P \leq 0.0001). For JNK inhibitor-treated larvae, decrease in CIFI for those fed 1, 5, and 10 µmol/L JNK-IN-8 of 2.7, 4.6, and 11.4 folds, respectively for GFP fluorescence (STT.P \leq 0.002) and of 3.4, 5.1, and 8.3 folds, respectively, for MMP1-related fluorescence (STT.P \leq 0.0005). For MMP1 inhibitor-treated larvae, decrease in CIFI for those fed on 5, 10, and 25 μ mol/L batimastat of 3.5, 4.8, and 8.4 folds, respectively, for GFP fluorescence (STT.P \leq 0.0002), and of 5.2, 5.9, and 10.3 folds, respectively, for MMP1-related fluorescence (STT.P = 0.02, 0.07, and 0.06, respectively).

 $75 \pm 5\%$ for MMP13 (Fig. 5). All differences were highly signif-318 icant (P < 0.001) with median duration of survival of 47, 32, and 319 52 months for MMP2, 9, and 13 positively stained tumors versus 320 228 months in all cases of negatively stained tumors. These 321 corresponded to relative risks (RR) of death of 9.04 (95% CI, 322 5.32-15.36), 4.69 (95% CI, 2.89-7.62), and 4.87 (95% CI, 2.98-323 7.97), respectively. Results for S100A4 with a cutoff of 5% were 324similar to that for individual MMPs with only $9 \pm 4\%$ surviving 325versus $80 \pm 4\%$ for unstained tumors, median survival time of 46 months versus 228 months ($\chi^2 = 71.8$, P < 0.001), and RR of 326 327 patient death of 9.96 (95% CI, 5.87-16.9; Supplementary Table 328 S3). Patients with tumors stained positively for all three MMPs 329showed no significant increase in mortality (7% \pm 6%), decrease 330

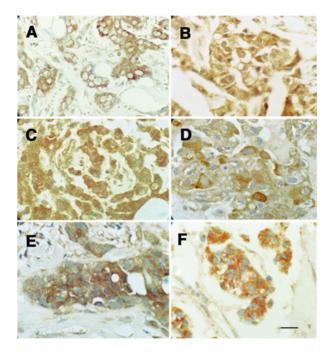


Figure 4.

Immunohistochemical staining of different breast carcinomas with antibody to MMP2 (**A**), MMP9 (**B**), or MMP13 (**C**) showing strong brown staining of the carcinoma cells' cytoplasm. Incubation with antibody to S100A4 (**D**) or S100P (**E**) showing strong, bead-like, cytoplasmic staining. **F**, Incubation with antibody to MMP2 with brown chromophore and to S100A4 with red chromophore showing most carcinoma cells were stained by both antibodies. Tumors were selected to show strong staining in **A** to **C** for their respective MMP, but the same tumor was stained in **D** to **F** as in **A** (×180; scale bar, 20 µm).

333 in median survival time (30 months), or increase in RR (4.96; 95% 334CI, 2.99-8.24) than staining for either MMP2 or MMP9 separately (Supplementary Table S2). When all three MMPs were included in 335 336 Cox's multivariate regression analysis (Materials and Methods), 337 the individual contributions made to the time of patient demise 338 showed that staining for MMP2 (P < 0.001) and that for MMP9 339 (P = 0.025) were independently significant while that for MMP13 340was not (Supplementary Table S3).

341 Association of MMPs with S100A4 and patient survival

342 Results for IHC staining for the 3 MMPs using a 5% cutoff were 343cross-tabulated against pathologic variables and IHC staining for S100A4, S100P (29), estrogen receptor α (ER α), progesterone 344 345receptor (PgR), c-erbB-2 (Her2), cytokeratin 5/6 (CK5/6), and CK14 (32). All these variables have been reported to influence 346347survival times in the same set of patients (26). Positive staining for 348 each of MMP2, 9, and 13 was associated strongly with positive staining for \$100A4 when using a 5% cutoff for \$100A4. This 349350 association was slightly reduced with staining for S100P using a 3515% cutoff (Table 2). Significance of association was much more marked for staining for S100A4 than for S100P when using a 1% 352cutoff. There was also a significant association with staining for 353 354CK5/6 and usually for CK14 (Table 2). Positive staining for any 355MMP was not significantly associated with involved lymph nodes, 356 high tumor grade, large tumor size, nor with positive staining for 357 ERα, PgR, or c-erbB-2 (Table 2). There was also a highly significant 358association of staining for each pair of MMPs (Table 2 and 359 Supplementary Table S4).

When staining for S100A4 was tested for its relative probability 361 of association (RA) with that for the three MMPs using binary 362 logistic regression, that with MMP2 was strongest at 4.21 (P < 363 0.001), that with MMP9 of 2.41 was not significant, and that with 364 MMP13 of 2.17 (P = 0.051) was very nearly significant. When 365 staining for each of the MMPs, in turn, was assessed with staining 366 for S100A4, CK14, ERa, PgR, and c-erb-2, only that for S100A4 367 and partially that for CK14 proved to be significant (Supplemen-368 tary Table S5). When repeated using a different cutoff for S100A4 369 (1% instead of 5%; Table 2) and additionally including that for 370 S100P, staining for MMP2 was most closely associated with that 371for S100A4 (Supplementary Table S5). To determine whether the 372 three MMPs were independent of \$100A4 when related to patient 373 survival, they were included in a series of Cox's multivariate 374regression analyses (Materials and Methods; Supplementary 375Table S3). When a single MMP and S100A4 were only included, 376 staining for S100A4 always emerged as the most significant 377 association with patient survival time. Similar results were 378 obtained if staining for S100A4 and all three MMPs were included 379 in the same analysis, S100A4 emerged as the most significant 380 association followed by MMP2 and then MMP9, whereas that due 381 to MMP13 was not significant (Supplementary Table S3). 382383

To determine whether there was coexpression of the MMPs and S100 proteins, two breast carcinomas were chosen that were either moderately or strongly stained for MMP2, and these were IHC restained for S100A4/P, 3 MMPs, CK5/6, and CK14. Exactly the same areas were examined for each antigen. The percentage of stained cells for \$100A4 was not significantly different from that for MMP2 and MMP13, while staining for S100P was not significantly different from that for MMP9 (Supplementary Fig. S6; and Supplementary Table S6). Staining for S100A4 or MMP2 was also not significantly different from that for CK5/6, but only in the MMP2 moderately-stained carcinomas; all the other paired combinations were significantly different (Supplementary Fig. S6; and Supplementary Table S6). When serial sections from three breast carcinomas strongly-staining for MMP2 were doubly IHC-stained for S100A4 with red and for MMP2 with brown chromophores on the same section, there were (mean \pm SE) 80.2 \pm 2.2% doubly stained cells, $6.9\% \pm 0.9\%$ cells stained red for S100A4, $2.9\% \pm$ 0.4% cells stained brown for MMP2 and 9.1% \pm 1.5% unstained cells (ANOVA, F = 669.3, 3 df, P < 0.001; Supplementary Fig. S7). Thus, S100A4 is associated with and partially confounded for patient survival by the three MMPs to varving degrees.

Discussion

We have shown for the first time that S100A4 can induce 405metastasis in the Drosophila model and that the oncogene Ras^{Val12} 406 largely fails in this respect. The increase in number of larvae 407 bearing VNC metastases (10-fold), in ASSM (24-fold), and in 408 CIFI (4.1-fold) for Ras^{Val12}/S100A4 over Ras^{Val12} larvae demon-409strates clearly that S100A4 promotes extensive dissemination 410 to the VNC, as well as elsewhere in the larvae (Supplementary 411 Fig. S1). The reason for the differences in fold increases is due to 412the method of measurement, the CIFI included GFP fluorescence 413due to the primary as well as the metastases, whereas the first 414 two parameters relate only to the metastases. That larvae contain-415ing Ras^{Val12} and inactive S100A4 $\Delta 2$ genes (20) show significantly 416 less metastases (Fig. 1A, 2A, and B), demonstrates that the 417migratory/invasive ability of S100A4 (20) is required for its 418 metastatic ability. That S100A4 larvae produce no tumors at all 419

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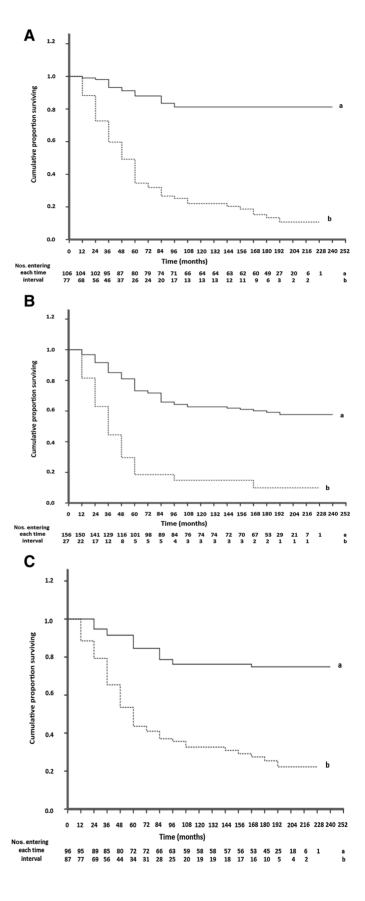


Figure 5.

Association of immunohistochemical staining for MMPs with overall time of patient survival. Cumulative proportion of surviving patients as a fraction of the total for each year after presentation for patients with carcinomas classified as negatively-stained (set a solid line) or positively-stained (set b, dotted line) is shown for MMP2 (**A**), MMP9 (**B**), and MMP13 (**C**). Numbers of patients entering each year are shown below. The two curves are highly significantly different in each case (Wilcoxon statistic $\chi^2 = 71.81$, 32.50, or 41.90 for **A**, **B**, or **C**, respectively, 1 df, P < 0.001). Further details are shown in Supplementary Materials.

Table 2. Association of IHC staining for MMPs with other tumor variables

			Statistical significance ^c	
Tumor variable ^a	Patient ^b no.	MMP2	MMP9	MMP13
Lymph nodes	139	0.271	0.564	0.681
Grade	164	0.997	0.656	0.156
Tumor size	177	0.467	0.937	0.985
MMP2	183	_	9.0×10^{-7}	1.7×10^{-12}
MMP9	183	9.0×10^{-7}	_	1.2×10^{-7}
MMP13	183	1.7×10^{-12}	1.2×10^{-7}	_
S100A4 (5%)	183	6.6×10^{-9}	2.9×10^{-4}	2.4×10^{-6}
S100A4 (1%)	183	O ^d	1.2×10^{-7}	1.9×10^{-8}
S100P (5%)	163	2.3×10^{-7}	2.2×10^{-3}	1.3×10^{-7}
S100P (1%)	163	1.6×10^{-4}	0.012	2.4×10^{-5}
CK14	172	5.7×10^{-7}	2.8×10^{-3}	0.372
CK5/6	173	1.6×10^{-6}	0.035	5.6×10^{-3}
ERα	181	1.00	1.0	1.0
PgR	172	0.995	0.983	0.549
C-erbB-2	183	0.660	1.00	0.809

^aLymph nodes with or without tumor deposits; grade, histologic grades I and II vs. grade III; tumor size <5 cm vs. >5 cm in diameter; presence or absence of IHC staining for molecular variables using 5% cutoff for MMP2, MMP9, MMP13, S100A4 (5%), S100P (5%), ERα, PgR, c-erbB-2, and using a 1% cutoff for S100A4 (1%), S100P (1%), CK14 and CK5/6

^bNumber of patients from original 183.

^cProbability P from Fisher exact test using the Holm-Bonferroni correction calculated as $1 - (1 - P)^n$, where n = 12 (Materials and Methods). ^dUncorrected $P = 7.7 \times 10^{-18}$

422 (Fig. 1A, 2A, and B) demonstrates that S100A4 alone is non-423oncogenic, consistent with previous results in our S100A4 transgenic mice (33). The increases in Ras and GFP proteins of 3.5- to 4-424 425fold (Table 1) are consistent with the increase in GFP fluorescence 426 of about 4-fold (Fig. 2B) and probably represent the increase in overall tumor mass between the Ras^{Val12} and the Ras^{Val12}/S100A4 427428larvae.

429 In agreement with different genetically manipulated Ras oncogenic systems in Drosophila (16, 18), the levels of endogenous 430activated phospho-JNK and MMP1 rise significantly in Ras^{Val12}/ 431S100A4 compared with Ras^{Val12}larvae (Table 1). The rise in MMP1 432protein is of the same order as the increase in fluorescently-labeled 433434antibodies to MMP1. That JNK is indeed a downstream effector of Ras^{Val12}/S100A4 for metastasis is demonstrated by the reduction 435436 in the number with metastases and their ASSM in female Bsk^{DN}expressing larvae compared to the male unsuppressed larvae (Fig. 4372C). That these suppressed values for Ras^{Val12}/S100A4 are not 438 significantly different from those of the *Ras^{Val12}* larvae (Fig. 2C) 439440 suggests that the predominant driver of the JNK-link to metastasis 441 is the overexpression of \$100A4. The 4.7-fold fall in the immuno-442 fluorescently-detectable β-galactosidase in the female, suppressed Ras^{Val12}/S100A4 larvae demonstrates that JNK needs to be acti-443vated to stimulate metastasis. Because of the level of JNK protein is 444relatively constant between Ras^{Val12} and Ras^{Val12}/S100A4 larvae 445446(Table 1), S100A4 probably triggers activation of JNK by stimulating its increase in phosphorylation (24). Results using 10 447 448 µmol/L JNK-IN-8 confirm that JNK-induced phosphorylation 449of c-Jun is a necessary step in the S100A4-triggered pathway for 450metastasis. That there is a fall in CIFI for immunofluorescently 451detectable MMP1 (Fig. 3B) positions JNK before MMP in any 452pathway (16). Moreover, the fact that the MMP1 inhibitor, batimastat (25) inhibits ASSM and CIFI for GFP in the Ras^{Val12}/ 453S100A4 larvae places MMP1 on the direct pathway to metastasis. 454455The order of this novel \$100A4-induced metastatic pathway 456is: S100A4->phospho-JNK->c-Jun->MMP1->metastasis. Thus, 457S100A4 appears to replicate the loss of function of suppressor genes scrib and lethal2 (17) or Her2 activation in the JNK/MMP 458pathway (16, 18). In transgenic mice or chemically transformed 459460rat mammary cells, S100A4 combines with oncogenic Neu (Her2;

ref. 6) or Ras (4), respectively to stimulate, via the cytoskeleton, cell migration, and then subsequent events for invasion/metastasis (34). However, the involvement of this novel pathway has hitherto been unreported.

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The relevance of our unique Drosophila model for S100A4 has 466 been pursued in human breast cancer. IHC staining of our cohort 467of 183 breast carcinomas for the individual MMPs2, 9, 13 468 demonstrates 15% to 47% primary tumors are stained positively 469 using a cut-off of 5%, in approximate agreement with previous 470 reports (35, 36). Here we show that the overall duration of 471survival of patients with positively-stained carcinomas is highly 472significantly worse than for those patients classified as not staining 473for one of MMP2, 9, or 13 (Fig. 5), in agreement with results 474for MMP2 in hepatocarcinoma (37), skin melanoma (38) and 475for MMP13 in breast (36) and colon cancer (39). In contrast, 476 MMP9 has been reported to be a favorable indicator in lymph-477node-negative breast cancer (40). This favorable prognosis may 478depend on the much higher cutoff employed, because of our 479node-negative group showed no significant difference (Wilcoxon 480 $\chi^2 = 2.63, 1 \text{ df}, P = 0.11$). This difference was significantly greater 481 for MMP9 staining in our node-positive patients ($\chi^2 = 18.40, 1 \text{ df}$, 482P < 0.001). The other two MMPs showed similar significant 483differences in node-negative and node-positive patients (MMP2 484 $\chi^2 = 25.46$ and 25.39; MMP13 $\chi^2 = 14.91$ and 12.93, respec-485tively). These results may suggest that MMP9 operates later than 486the other two MMPs at a post lymph-node-spreading stage in the 487 disease process. 488

Overall, the RR of patient death in separate univariate analyses 489 is greatest for patients with tumors stained for S100A4 (9.96), 490followed closely by those stained for MMP2 (9.04), then for 491 MMP13 (4.87), and finally for MMP9 (4.69; Supplementary Table 492S3). However, the antibodies used here to detect the MMPs do not 493discriminate between inactive precursors or cleaved active MMPs 494 and do not detect inhibitory TIMPs (41). Usually in cultured cells, 495S100A4 increases expression of MMP precursors and this results 496in an enhanced proteolytic activity and cell invasion/metastasis 497(42, 43). Moreover, S100A4 can act both intracellularly (43, 44) 498and extracellularly via RAGE receptors (45, 46) to stimulate MMP 499production. The fact that *Bsk*^{DN} inhibits \$100A4-induced MMP1 500 503and metastasis to the VNC in our Drosophila model (Fig. 2C and D) 504suggests that MMP1 is produced by the tumor cells and not by reactive stromal cells (47), consistent with immunohistochemical 505506results in our human breast cancers. In contrast to the Drosophila 507model, the three JNK proteins in human cancers can exert both 508 pro- and anti-oncogenic effects depending on the cell type and 509 cross-talk with other kinases (48, 49). Thus, the oncogenic effect 510of activated INK cannot be determined in human cancers from the 511measurement of its level alone, and hence was not attempted here.

512Upon manipulation in cultured cells, S100A4 has been 513reported to control production of a single MMP, one of MMP2, 5149, or 13, depending on the source and sometimes the report (42-45). In contrast, we show here that positive staining for each 515516MMP2, 9, or 13 is separately and in combination very strongly 517associated with S100A4 and to a lesser extent with S100P (Table 2). The significant association of staining for MMP2, 9 with the 518basal cell markers CK5/6, CK14 has been reported previously 519520(50), predominantly placing these MMPs, together with S100A4 521and S100P, in the most aggressive subtype of breast cancers (26). 522When tested for RA of staining for S100A4 with the other three 523MMPs together, S100A4 is more likely to occur with MMP2, and 524the higher significant RA of MMP2 for S100A4 over a combination 525of other proteins confirms this result (Supplementary Table S6). 526Thus, S100A4 is more associated with MMP2, 13, and S100P more 527with MMP9, at least at the cellular level (Supplementary Table S6). 528This differential association in the tumor raises the novel possi-529bility of synergistic interactions between the S100 proteins (29) 530occurring via different target MMPs.

531Multiple longitudinal comparisons with survival time for all 532three MMPs together in multivariate analysis shows that only 533MMP2 and MMP9 are independently significant, whereas the contribution of MMP13 is confounded by that due to the other 534535two MMPs (Supplementary Table S3). These results suggest partial 536overlap occurs between MMP2/MMP9-related pathways and 537MMP13-related pathways, whereas those related to MMP2 and MMP9 are more separate. This result is consistent with their 538539function, MMP13 is a collagenase which is required to cut col-540lagen fibrils first, before the two gelatinases, MMP2 or MMP9, can 541digest the remainder (51). When S100A4 and each MMP are tested 542in combination, the order of reduction in RR for \$100A4 is MMP2 543(42% reduction), then MMP13 (27% reduction) and finally 544MMP9 (11% reduction), whereas the reduction in RR for each 545MMP separately with S100A4 is similar (44%, 40%, and 43%,

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respectively; Supplementary Table S3). These results suggest the 547pathways that S100A4 may trigger leading to premature death 548from metastatic disease overlap, to some extent, with those 549triggered by the three MMPs, the most overlap being with 550MMP2-related and then with MMP13-related pathways. The 551results for the close association of S100A4 and MMP2 are con-552firmed at the level of the cell where 91% of \$100A4-containing 553cells also contain MMP2 and 96% of MMP2-containing cells also 554contain \$100A4 (Supplementary Fig. \$7). The considerable 555enhancing effect of \$100P on \$100A4-linked patient demise 556(29) may then be attributable, at least in part, to \$100P targeting 557different MMPs from those targeted by S100A4 (Supplementary 558559Table S6). This differential targeting of MMPs by S100 proteins is a novel mechanism for generation of the known synergy between 560different MIPs in the development of many cancers. 561

Disclosu	re of	Pote	ential	Conflicts	of Interest	
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Q8 563 No potential conflicts of interest were disclosed.

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Rudland	567
Acquisition of data (provided animals, acquired and managed patients,	568
provided facilities, etc.): T.M. Ismail, D. Bennett, P.S. Rudland	569
Analysis and interpretation of data (e.g., statistical analysis, biostatistics,	570
computational analysis): T.M. Ismail, D. Bennett, A.M. Platt-Higgins,	571
M. Al-Medhity, P.S. Rudland	572
Writing, review, and/or revision of the manuscript: T.M. Ismail, D. Bennett,	573
A.M. Platt-Higgins, M. Al-Medhity, R. Barraclough, P.S. Rudland	574
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data, constructing databases): T.M. Ismail, A.M. Platt-Higgins, P.S. Rudland	576
Study supervision: T.M. Ismail, P.S. Rudland	Q9 577

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