Dystrophin Is a Tumor Suppressor in Human Cancers with Myogenic Programs

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Abstract

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NCBI Gene Expression Omnibus (GEO): GSE53021, for genome-wide 250K SNP genotypes of 40 myogenic-cancers and normal controls.

AUTHOR CONTRIBUTIONS
Many common human mesenchymal tumors, including gastrointestinal stromal tumor (GIST), rhabdomyosarcoma (RMS), and leiomyosarcoma (LMS), feature myogenic differentiation.\(^1\)–\(^3\) Here we report that intragenic deletion of the dystrophin-encoding and muscular dystrophy-associated \textit{DMD} gene is a frequent mechanism by which myogenic tumors progress to high-grade, lethal sarcomas. Dystrophin is expressed in nonneoplastic and benign counterparts for GIST, RMS and LMS, and the \textit{DMD} deletions inactivate larger dystrophin isoforms, including 427kDa dystrophin, while preserving expression of an essential 71kDa isoform. Dystrophin inhibits myogenic sarcoma cell migration, invasion, anchorage independence, and invadopodia formation, and dystrophin inactivation was found in 96%, 100%, and 62% of metastatic GIST, embryonal RMS, and LMS, respectively. These findings validate dystrophin as a tumor suppressor and likely anti-metastatic factor, suggesting that therapies in development for muscular dystrophies may also have relevance in treatment of cancer.

Human cancers featuring myogenic differentiation include LMS, RMS, and GIST. GIST – although most closely resembling interstitial cells of Cajal (ICC) – often express myogenic differentiation markers, such as smooth muscle actin and calponin\(^4\)–\(^7\). Presumptive initiating mutations have been identified in these myogenic cancers, including germline \textit{TP53} mutations in patients with LMS and RMS\(^8,9\) and gain-of-function \textit{KIT} or \textit{PDGFRA} mutations in patients with GIST\(^10\)–\(^12\). Somatic mutations contribute to tumorigenic progression in myogenic cancers, e.g. cell cycle dysregulation by \textit{CDKN2A} or \textit{TP53} inactivation in GIST\(^13,14\), but few of these genetic progression mechanisms in myogenic cancers have been characterized.

To identify shared tumorigenic mechanisms in myogenic cancers, we performed genome-wide Affymetrix 250K single-nucleotide polymorphism (SNP) assays. These studies revealed intragenic deletions in the Duchenne and Becker muscular dystrophy gene, \textit{DMD}\(^15\) in 25 of 40 high-grade myogenic cancers (63%), including 19 of 29 GISTs, 3 of 4 RMS, and 3 of 7 LMS (Fig. 1a). Although \textit{DMD} is an X-linked gene, the deletions were found in both male and female patients, including 9 of 13 (69%) GISTs in men and 10 of 16 (63%) GISTs in women (Supplementary Table 1). \textit{DMD} deletions in myogenic cancers were not present in companion non-neoplastic tissues, attesting to somatic origin (Fig. 1b). \textit{DMD} deletions, when identified within a primary GIST, were perpetuated in subsequent metastatic lesions (Supplementary Fig. 1), and when identified in any GIST metastasis, were present in other metastases from the same patient (Supplementary Fig. 2). \textit{DMD} intragenic deletions were not detected in 58 non-myogenic sarcomas (Supplementary Table 2) and were observed only infrequently (4.3%) in 905 non-sarcoma human cancer cell lines in the Cancer Cell Line Encyclopedia program\(^16\), (Supplementary Fig. 3). These data show that the frequency of \textit{DMD} deletions is higher in myogenic cancers compared to non-myogenic tumors (\(P < 0.0001\)).

Whereas all \textit{DMD} deletions in cancers from men were nullizygous, the deletions in cancers from women were either nullizygous (\(n = 9\)) or heterozygous (\(n = 4\)) (Fig. 2a and Supplementary Table 1). Fluorescence \textit{in situ} hybridization for \textit{DMD} and the Xist inactive X chromosome marker\(^17\) showed that heterozygous \textit{DMD} deletions targeted the active X
chromosome (Fig. 2b). Therefore, both nullizygous and heterozygous DMD deletions in female patients caused complete DMD inactivation.

DMD is the longest known human gene, composed of 79 coding exons spanning 2.2 megabases of the genome, with various transcriptional start sites. Multiplex ligation-dependent probe amplification (MLPA) copy number assessment for each of the DMD coding exons revealed intragenic DMD deletions in 24 of 56 high-grade myogenic cancers (43%) (Fig. 3, Supplementary Fig. 4 and Supplementary Table 3), all of which were predicted to abrogate expression of the largest dystrophin isoform (427kDa), encoded by DMD exons 1–79. By contrast, intragenic DMD deletions were not found in 20 benign tumor counterparts for GIST, RMS and LMS (11 low-risk GISTs, 2 rhabdomyomas and 7 leiomyomas), despite high levels of dystrophin expression (Supplementary Table 3). The dystrophin 427kDa isoform was expressed strongly in normal tissue and benign counterparts for GIST, RMS and LMS (Fig. 4), but was undetectable or weakly expressed in 96% of metastatic GISTs (26 of 27), irrespective of whether they contained KIT or PDGFRA mutations (Fig. 4b and Supplementary Table 4). Similarly, dystrophin 427kDa expression was undetectable or weak in 100% of metastatic embryonal RMS (eRMS) (9 of 9), and 62% of metastatic LMS (8 of 13) (Fig. 4c,d and Supplementary Table 4). Dystrophin 427kDa was also downregulated in 75% of primary “high-risk” GISTs (i.e., GISTs having histologic criteria predictive of metastasis), consistent with the SNP evidence that DMD dysregulation is positively selected for in clinically-aggressive primary tumors, even prior to metastasis (Fig. 4b). Expression of 427kDa dystrophin was not detected in 46 non-myogenic sarcomas (Supplementary Table 5). By contrast, expression of dystrophin isoform Dp71 (71kDa), encoded by exons 63–79, is preserved in cancers with DMD deletions (Fig. 5). Dp71 is also expressed in non-myogenic sarcomas (Supplementary Fig. 5), in keeping with reports that Dp71 expression is ubiquitous in nonneoplastic tissues, other than skeletal muscle.

Dystrophin biologic function was evaluated by re-expressing dystrophin in DMD-inactivated GIST, RMS and LMS. Re-expression of dystrophin 427kDa is challenging, due to the large size of the cDNA construct. Therefore, we used a miniDMD construct lacking exons 17–48 which encode a spectrin-like domain (Fig. 3). This miniDMD construct is biologically relevant, inasmuch as it restores crucial aspects of dystrophin function in gene therapy for muscular dystrophy patients. MiniDMD transfection into DMD-inactivated GIST, eRMS and LMS induced 240kDa dystrophin expression at levels that are physiologic for dystrophin 427kDa expression in low-risk (indolent) GIST, skeletal muscle and myometrium, respectively (Supplementary Fig. 7). Dystrophin re-expression inhibited invasiveness and migration in GIST, eRMS and LMS, but showed no effect in the non-myogenic fibrosarcoma cell line HT-1080 (Fig. 6a,b), and reduced the number of viable cells in eRMS and LMS, but not in GIST, fibrosarcoma, Ewing’s sarcoma or HEK 293 cells (Supplementary Fig. 8). Dystrophin re-expression inhibited anchorage-independent growth in DMD-inactivated GIST, eRMS and LMS (Fig. 7a), but not in comparator non-myogenic sarcomas.
fibrosarcoma or Ewing’s sarcoma (Fig. 7a and Supplementary Fig. 9). These studies support the hypothesis that DMD inactivation enhances metastatic potential selectively in cancers with myogenic programs.

Dystrophin provides a structural link between the actin-based cytoskeleton and extracellular matrix\(^{22,23}\), which is consistent with our evidence that dystrophin regulates invasion and migration in myogenic cancers. Metastases account for 90% of cancer-related deaths\(^{24,25}\), and some invasive cancer cells feature invadopodia, which are actin-rich membrane protrusions regulating metastatic behavior\(^{26}\). Restoration of DMD expression inhibited invadopodia formation in GIST and LMS (Fig. 7b), also consistent with dystrophin function as an anti-metastatic factor.

The genomic, clinicopathological and functional evidence herein demonstrate dystrophin tumor suppressor roles contributing to permissiveness for metastatic behavior in human cancers. The relevance of these findings is supported by reports of spontaneous RMS in DMD-inactivated mdx mice\(^{27-29}\). Similarly, reports of RMS in Duchenne muscular dystrophy patients suggest that germline DMD inactivation may predispose to myogenic cancer\(^{30,31}\). DMD is a large gene, but of the >40 human genes that are more than 1 megabase in genomic length, only DMD is known to have frequent deletions that seem subject to strong selective pressure in pre-metastatic GIST. Several observations support DMD inactivation as a driver event in myogenic cancers: 1) frequent DMD deletions are not found in non-myogenic cancers (Supplementary Fig. 3), most of which have more complex genomic landscapes than those in RMS and GIST\(^ {32}\), indicating that clonal DMD deletions in advanced myogenic cancers are not attributable merely to cytogenomic complexity; 2) DMD deletions are not found in benign precursors to GIST (Fig. 4b) but rather are remarkably frequent late events in GIST progression and are found in subclones which – based on the genomic evidence – overgrow DMD-wildtype subclones in the same tumors; 3) although a late event in tumorigenesis, the same DMD deletion found in one metastasis can be detected in all other metastases from the patient (Supplementary Fig. 2), consistent with a biologic advantage for dystrophin inactivation. These genomic lines of evidence, coupled with demonstration that DMD-restoration is only impactful in myogenic cancers (Fig. 6,7), suggests that DMD mutations in myogenic cancers are classic tumor suppressor events.

It is unclear whether the DMD mutation rate is particularly high in myogenic cancers, perhaps due to mechanisms such as transcription-associated recombination\(^{33}\), but in any case the combined evidence summarized above suggests that DMD inactivation may be more than simply a passenger event in myogenic cancer. Previously, DMD deletions were demonstrated in 5.5% of malignant melanoma\(^ {34}\), but it is unknown whether those deletions had functional consequences. In clinically-advanced myogenic cancers, we show that DMD inactivation abolishes the dystrophin 427kDa expression found in normal tissue and benign counterparts for GIST, LMS and RMS, while preserving dystrophin 71kDa protein, which appears to be an obligate factor in these cancers. Dystrophin interacts with a complex of sarcolemmal proteins and glycoproteins known as dystrophin-associated proteins\(^{22}\), and our demonstration of dystrophin tumor suppressor roles anticipates that other proteins in this complex might regulate tumorigenic functions. As one example, the dystrophin-related protein, utrophin, is a potential tumor suppressor in non-myogenic malignancies\(^{35}\), and
pharmacologic induction of utrophin overexpression in mdx mice prevents development of muscular dystrophy, suggesting that utrophin can compensate for dystrophin deficiency. Other treatment options to correct dystrophin defects are undergoing evaluation in clinical trials for Duchenne muscular dystrophy, and these approaches warrant evaluation as potential therapeutic agents in oncology. Identification of patients whose cancers have dystrophin defects might be expedited by immunohistochemical screening for loss of dystrophin 427kDa expression. Immunohistochemical assays show robust dystrophin expression in nonneoplastic myogenic cells (skeletal, cardiac, and smooth muscle) and in leiomyoma (benign smooth muscle tumors) (Supplementary Fig. 10a,c), whereas LMS (malignant smooth muscle tumors) often feature complete loss of demonstrable dystrophin expression (Supplementary Fig. 10b,c). Therefore, dystrophin dysregulation warrants evaluation as a prognostic factor in myogenic cancers, and as a potential point of therapeutic attack.

METHODS

Tumor and tissue samples

Snap-frozen tumor biopsies and matched normal tissue samples were from patients at Brigham and Women’s Hospital and Memorial Sloan Kettering Cancer Center. All samples were collected with institutional review board approval.

SNP arrays

High molecular weight genomic DNA was isolated using QIAamp DNA Mini Kit (QIAGEN) and analyzed by Affymetrix 250K SNP array. DNA was digested with Nsp1, and linkers were ligated to the restriction fragments to permit PCR amplification. The PCR products were purified and fragmented by treatment with DNase I, then labeled and hybridized to microarray chips. The positions and intensities of the fluorescence emissions were analyzed using dChip software. Array intensity was normalized to the array with median intensity. Median smoothing was used to infer copy number.

Fluorescence in situ hybridization and Xist evaluation

BACs RP11-42E12 (DMD, chromosome Xp21.1; start position 32,642,966 GRCh37/hg19; end position 32,848,014; targeting DMD introns 4–9) and RP11-939O17 (DMD, chromosome Xp21.1; start position 32,844,043; end position 33,018,177; targeting DMD introns 2–4) were obtained from BAC/PAC Resources (Children’s Hospital, Oakland, CA). BAC DNA was labeled using a nick translation kit with Spectrum Orange-11-dUTP (DMD). As a control for X chromosome copy number a centromere X probe (DXZ1, spectrum aqua probe) was obtained from Abbott Laboratories (Vysis, Abbott Park, Illinois). A fosmid clone overlapping the Xist locus (G248P8779H11; Xq13.2; start position 73,038,817; end position 73,075,707) was obtained from BAC/PAC Resources (Children’s Hospital, Oakland, CA). The fosmid DNA was labeled by nick translation with Spectrum Green-11-dUTP, and used in a triple-hybridization with the probes for DMD (spectrum orange) and centromere X (aqua) according to standard protocols.
Multiplex Ligation-dependent Probe Amplification (MLPA)

The MLPA procedure and capillary electrophoresis were performed using SALSA MLPA kits P034-A2 and P035-A2 from MRC-Holland. The combined P034 and P035 kits contain probes for each of the 79 DMD coding exons and for the alternative exon 1 Dp427c. MLPA reactions were conducted using a G-Storm GS1 thermal cycler (Gene Technologies Ltd) with fragment analysis by ABI-3130XL Genetic Analyzer and GeneMapper software (Applied Biosystems). Raw data were received as peak heights, as a measure of peak intensity, for each of the probes.

Purification of murine interstitial cells of Cajal (ICC)

Murine ICC (~500,000) were isolated as Kit+CD44+CD34− cells from the hematopoietic marker-negative (CD45−F4/80−CD11b−) fraction of the gastric tunica muscularis of day 7–15 BALB/c mice (n = 52, in 5 cohorts) as described previously41,42. The fidelity of ICC sorting was validated by demonstrating a >2.5-fold increase in KIT protein expression in sorted compared to unsorted cells and by lack of expression of the smooth muscle marker Myh11 and the pan-neural marker PGP 9.5 by western blotting.

Western blotting

Frozen tumor samples were diced in ice-cold lysis buffer (1% NP-40, 50 mM Tris-HCl pH 8.0, 100 mM sodium fluoride, 30 mM sodium pyrophosphate, 2 mM sodium molybdate, 5 mM EDTA, 2 mM sodium orthovanadate) on dry ice and homogenized with a Tissue Tearor Homogenizer for 3 seconds, 3–5 times, on ice, and the cell lysate was then rocked overnight at 4°C. Lysates were cleared by centrifugation at 14,000 rpm for 30 min at 4°C, and lysate protein concentrations were determined using a Bio-Rad protein assay (Bio-Rad Laboratories Hercules, CA, USA). Electrophoresis and western blotting were performed using standard techniques. The hybridization signals were detected by chemiluminescence (Immobilon Western, Millipore Corporation, MA) and captured using a FUJI LAS1000-plus chemiluminescence imaging system (Fuji Film, Tokyo, Japan). Primary antibodies were DYS1 (Novocastra, NCL-DYS1, raised against the dystrophin rod domain, amino acids 1181 and 1388, detects 427 kDa dystrophin isoform), DYS2 (Novocastra, NCL-DYS2, raised against the C-terminal 17 amino acids of dystrophin, detects 240 kDa mini-dystrophin), 7A10 (Santa Cruz, sc-47760, raised against amino acids 3200–3684 of dystrophin, detects Dp71), and GAPDH (Sigma, G8795).

Cell lines

GIST-T1 was generously provided by Dr. Takahiro Taguchi43. HT-1080 and HEK 293 were obtained from the American Type Culture Collection. All other cell lines were developed in the Jonathan Fletcher laboratory at Brigham and Women’s Hospital. GIST-T1 and GIST430 were established from metastatic GISTs with DMD exon 1 and exons 1–9 deletions, respectively. RMS176 was established from a metastatic eRMS with DMD exons 1–7 and exons 21–44 deletions. LMS04 was established from a metastatic LMS without apparent DMD deletion but with complete loss of 427kDa dystrophin expression. The HT-1080 fibrosarcoma44 and EWS502 Ewing’s sarcoma were non-myogenic controls with wildtype DMD.
**Stable transfections**

GIST-T1, GIST430, RMS176, LMS04 and EWS502 cell monolayers were disaggregated with trypsin and resuspended in Amaza Nucleofector solution V (Amaza Biosystems) at a concentration of 1 x 10^6 cells per 100 μl. Nucleofection was performed using program T-030 on a Nucleofector II machine (Amaza Biosystems). One microgram of pCR3.1-EGFP or pCR3.1-miniDMD plasmid was used for electroporation. Transfected cells were selected with G418 for 5 days before analyses.

**Dp71 siRNA knockdown**

Dp71-specific siRNA, targeting the unique region in exon 1 of Dp71, was obtained from Invitrogen (sequence provided in Supplementary Table 6). Control siRNA was obtained from Invitrogen (Stealth RNAi™ siRNA negative control medium GC Duplex, Catalog # 12935-300). siRNA was delivered into RMS176 and RMS843 cells by nucleofection (Amaza Biosystems) as described previously.

**Cell viability assays**

Viability studies were performed using the CellTiter-Glo luminescent assay (Promega, Madison, WI). Cells were plated at 2,000 cells per well in a 96-well flat-bottomed plate (Falcon, Lincoln, NJ). Luminescence was analyzed using a Veritas microplate luminometer (Turner Biosystems, Sunnyvale, CA).

**Soft agar assay**

Cells were plated in 35mm dishes after stable transfection of EGFP or miniDMD. The cells were incubated for 3–4 weeks and then stained with 1 ml of 1 mg/ml methyl thiazol tetrazolium (MTT) for 3 hours. Colonies were counted manually. All experiments were performed in triplicate.

**Quantitative cell invasion and migration assays**

0.3 ml serum free media containing 0.3×10^5 GIST-T1, GIST430, RMS176 or LMS04 cells were plated for invasion assays in BD BioCoat™ Matrigel Invasion Chambers (BD Biosciences, Catalog # 354480) and for migration assays in BD BioCoat™ 8.0 μm PET Membrane 24-well Cell Culture Inserts (BD Biosciences, Catalog # 354578). The wells were fed with 0.5ml RPMI Media 1640 (Invitrogen) containing 15% FBS and incubated in a humidified incubator, at 37°C, 5% CO₂ for 144 hours (GIST-T1), 72 hours (GIST430) or 24 hours (RMS176, LMS04, and HT-1080). The media from the inside of the insert was aspirated, and the interiors of the inserts were gently swabbed to remove non-invasive or non-migratory cells. Inserts were transferred to new wells containing 400 μl Cell Stain Solution (Cell Biolabs, Inc) and incubated for 10 minutes at room temperature, then rinsed two times in a beaker of water. Then, the inserts were air dried and transferred to new wells containing 200 μl Extraction Solution (Cell Biolabs, Inc) and quantified at OD 560nm in a microplate reader.
Radius 2-D cell migration assay

Impact of dystrophin restoration on cell migration was determined using a 2-dimensional gap closure radius 24-well migration assay, according to the manufacturer’s instructions (Cell Biolabs, Inc).

Immunofluorescence

5 ×10^4 cells were plated on 10-mm glass coverslips coated with 1 μg/cm² fibronectin (Sigma-Aldrich). Cells were fixed in 4% paraformaldehyde (Electron Microscopy Sciences) for 15 min, permeabilized for 10 min. with 0.3% Triton X-100 (Thermo Fisher Scientific) for 10 min, incubated with primary antibodies overnight at 4°C, then with secondary antibody for 1 hour, and mounted on slides with FluorSave™ reagent (EMD Millipore Chemicals). Cells were stained for cortactin and MMP14 to identify invadopodia.

Immunohistochemistry

Immunohistochemistry was performed on tissue and tumor sections using DYS1 mouse monoclonal antibody (Novocastra, NCL-DYS1). Four micron slides were deparaffinized in xylene and hydrated in a graded series of alcohol. The deparaffinized slides were then boiled by microwave for 12 minutes in citrate buffer (pH 6). The IHC reactions were visualized by diaminobenzidine staining, using an EnVision+ system (Dako, Carpinteria, CA, USA).

Statistical analysis

Two-tailed unpaired t-tests were performed for comparison of means analysis. For 2×2 contingency tables, two-tailed P values were calculated using Fisher’s exact test.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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References


Figure 1.
Identification of somatic intragenic DMD deletions in human myogenic cancers. (a) dChip SNP log2 ratio copy number evaluations demonstrate intragenic DMD deletions in 25 of 40 (63%) primary or metastatic myogenic cancers. M denotes male and F denotes female (full clinicopathological details are provided in Supplementary Table 1). Panel on right depicts representative SNP profile with DMD deletion in GIST from a male patient. (b) SNP evaluations in matched cancer and non-neoplastic cell DNAs from the same patients, demonstrating tumor-restricted nature of DMD deletions. (c and d) DMD deletions in myogenic cancers from women. (c) dChip SNP analyses showing normal DMD copy number (N, case 28), nullizygous DMD deletion (Null, case 44) and heterozygous DMD deletions (cases 40 and 57 are metastatic GISTs; case 116-4 is metastatic LMS). (d) FISH and Xist analysis in cases 40, 57, and 116-4, showing DMD deletion in the Xist-negative
active X chromosome. Probes are for DMD (red), X centromere (blue), and Xist (green). Scale bars, 2 μm.
Figure 2.
MLPA evaluations of *DMD* exons 1–79 show intragenic deletions in 24 myogenic cancers.
Figure 3.
Loss of dystrophin 427kDa expression in most metastatic GIST, RMS and LMS. Western blotting with DYS1 demonstrates dystrophin expression in the normal tissue and benign tumor counterparts for GIST, RMS and LMS (a to d); low-risk GIST is a clinically indolent precursor to malignant GIST. Loss of dystrophin 427kDa expression is demonstrated in most metastatic GIST (b), RMS (c) and LMS (d). Patient 116 samples (d) are five successive LMS metastases, resected during an 8 year interval in the same patient.
Figure 4.
Expression of Dp71 dystrophin in myogenic tumors. Western blotting with dystrophin antibody 7A10 demonstrates Dp71 expression in primary GIST, metastatic GIST, and RMS cases, including those with loss of 427kDa dystrophin.
Figure 5.
Restoration of dystrophin expression inhibits invasiveness and migration in DMD-inactivated GIST, RMS and LMS, but not in a comparator non-myogenic fibrosarcoma (HT-1080). Three biological replicates of each experiment were evaluated; the error bars on the histograms indicate standard deviation of the mean. Scale bars, 100 μm. (a) miniDMD restoration in GIST, RMS and LMS inhibits invasion in Matrigel assays. GIST, RMS, LMS and fibrosarcoma cells were seeded and invasion toward 15% FBS was measured after 144 hours (GIST) or 24 hours (RMS, LMS and fibrosarcoma). Cells invading to the bottom of the membrane were stained and quantified at OD 560nm after extraction. (b) miniDMD restoration in GIST, RMS and LMS, but not in HT-1080 fibrosarcoma, inhibits migration, as assessed by Radius 2-D cell migration assay and by a complementary assay of migration toward 15% FBS on a polycarbonate membrane: migration was quantified at OD 560nm after extraction.
Figure 6.
Restoration of dystrophin expression inhibits anchorage-independent growth and invadopodia formation in DMD-inactivated myogenic cancers. Three biological replicates of each experiment were evaluated. (a) Stable miniDMD expression suppresses anchorage-independent growth in myogenic cancers (GIST, RMS and LMS) but not in non-myogenic comparators (fibrosarcoma HT-1080, Ewing’s sarcoma EWS502, and HEK 293) ($P<0.001$). Representative plates and mean colony numbers are shown (± standard error of the mean). (b) miniDMD restoration disrupts invadopodia formation in GIST and LMS cells. Top: Invadopodia are dot-like structures, staining with cortactin (blue) and MMP14 (red). Scale bars, 10 μm. Bottom: Percentages of cells with invadopodia in control (EGFP vector) vs. dystrophin-restoration conditions. The error bars on the histograms indicate standard deviation of the mean.