Laminin-1-induced migration of multiple myeloma cells involves the high-affinity 67 kD laminin receptor

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Summary The 67 kD laminin receptor (67LR) binds laminin-1 (LN), major component of the basement membrane, with high affinity. In this study, we demonstrated that human multiple myeloma cell lines (HMCL) and murine 5T2MM cells express 67LR. CD38bright+ plasma cells in the murine 5T2MM cells. 67LR has been shown to mediate the actions of LN through binding to CDPGYIGSR, a 9 amino acid sequence from the B1 chain of LN. MM cell migration was partially blocked by peptide 11, a synthetic nonapeptide derived from this amino sequence and also by a blocking antiserum against 67LR. Co-injection of peptide 11 with 5T2MM cells in a murine in vivo model of MM resulted in a decreased homing of 5T2MM cells to the BM compartment. In conclusion, LN acts as a chemoattractant for MM cells by interaction with 67LR. This interaction might be important during extravasation of circulating MM cells. © 2001 Cancer Research Campaign

Keywords: multiple myeloma; homing; laminin-1; 67LR; migration; extravasation

Multiple myeloma (MM) represents a B cell malignancy, characterised by a monoclonal proliferation of malignant plasma cells (Ludwig et al, 1999). During the main course of disease evolution, terminally differentiated B cells preferentially accumulate in the bone marrow (BM) and represent the main cell type. However, several studies indicated that the tumour clone also includes circulating B cells that might represent the ‘myeloma stem cell’. Clear evidence supporting this statement was provided by molecular techniques showing that MM patients have B cells in circulation harbouring the same immunoglobulin (Ig) gene rearrangements as the malignant BM plasma cells (Van Riet et al, 1989). Moreover it was demonstrated that these circulating MM cells show in their Ig genes the presence of somatic mutations without intraclonal variation (Bakkus et al, 1992). This implicates that MM originates from a B cell that has been antigen-selected in the germinal centre of the lymph node. The restricted localisation of MM cells in the BM during the initial stage of disease could be explained by a selective initial homing of circulating MM cells to the BM microenvironment and/or by the presence of a unique combination of survival and growth factors provided by the BM microenvironment not present in other tissues (Van Riet et al, 1998; Vanderkerken et al, 2000).

In analogy to the migration mechanisms used by normal leukocytes and metastatic carcinoma cells (Liotta, 1986; Butcher and Picker, 1996), the migration of MM cells to the BM is likely to be mediated by similar mechanisms. During lymphocyte homing and tumour cell invasion, cells attach to and degrade components of the extracellular matrix (ECM) before passing through. Previous studies already revealed the importance of interactions of malignant cells with the basement membrane during invasion. The basement membrane represents a specialised form of ECM, which separates the endothelium from the underlying interstitial stroma. It is composed of several components including collagen IV, glycoproteins such as laminin, fibronectin and proteoglycans (Vracko, 1974). Components of the ECM, including laminin, fibronectin and type IV collagen, as well as their proteolytic digestion products have been shown to stimulate the in vitro migration of a variety of normal and malignant cell types (Klominek et al, 1993). Laminin-1 (LN), a well characterised cell adhesion protein, represents the major non-collagenous glycoprotein of the basement membrane, involved in multiple important biological activities such as cell adhesion, migration, spreading, proliferation, growth and differentiation (Malinda and Kleinman, 1996). Very recently, Spessotto et al (2001) have demonstrated that different LN isoforms may evoke diverse cellular responses in different neoplastic lymphocytes. Invasion of the subendothelial basement membrane is mediated in part by interactions of tumour cells with laminin via specific cell surface receptors (Ramos et al, 1991). The 67 kD laminin-binding protein (67LR) represents a non-integrin-binding protein which binds LN with high affinity (Malinoff and Wicha, 1983). Expression of 67LR has been shown to be increased in neoplastic cells as compared to their normal counterparts and directly correlates with an enhanced invasive and metastatic potential (D’Errico et al, 1991; Kim et al, 1998). Additionally, suppression of the expression of the precursor of 67LR (37LBP) with antisense 37LBP/p40 RNA using a retroviral vector reduces the capability of lung cancer cell proliferation in vitro and tumour formation in vivo (Satoh et al, 1999), clearly indicating the role of 67LR in tumorigenicity.

In the present study, we demonstrated that MM cells express the 67 kD high-affinity LN receptor. In addition, we found that LN acts as a chemoattractant for MM cells by binding to 67LR.
MATERIALS AND METHODS

Human MM cell lines (HMCL)

Three well characterised HMCL (MM5.1, MMS.1 and U266) were selected for our experiments. They were cultured as described (Nilson, 1971; Okuno et al., 1991; Van Riet et al., 1997).

Patient samples

BM samples from 5 MM patients (pts) (age 52–94, mean 66.5; M/F, 3/5) were collected during standard diagnostic procedures. The study was approved by the local ethical committee. According to Durie and Salmon’s clinical staging system (1975), patients were distributed as follows: stage I, 1 pt; stage II, 2 pts; stage III, 2 pts. BM aspirates were collected in a heparinised syringe. Mononuclear cells were separated by density gradient centrifugation over Ficoll-Hypaque (Life Technologies).

Primary MM cells expressing the CD138 antigen were immunomagnetically separated using the Magnetic Cell Sorting System (MACS) (Miltenyi Biotech). Cells were incubated for 15 minutes at 4°C with MACS microbeads conjugated to a monoclonal mouse CD138 (syndecan-1) antibody (clone B-B4, isotype mouse IgG1) (40 μl 10⁷ cells). Cells were washed once in PBS supplemented with human albumin (4%), resuspended and separated on a column placed in the magnetic field of the MACS separator. CD138+ cells were retained in the column and eluted as positively selected cell fraction after removal of the column from the magnetic field. Cells were counted and viability was assessed with trypan blue. MACS purification produced a ≥ 97% pure preparation.

5T2 murine MM model

Male C57B1/KaLwRijHsd mice, 6–10 weeks old were purchased from Harlan CPB. Mice were housed under conventional conditions and fed ad libitum (license number LA 1230281). 5T2MM cells originated spontaneously in aging mice (Radl et al., 1979). The 5T2MM model displays characteristics analogous to human MM (Asosingh et al., 2000). Cells were prepared as described previously (Vanderkerken et al., 1997). Briefly, cells were obtained by flushing the BM of hind legs. Cells were purified by gradient centrifugation on lymphocyte M (Cedarlane), washed and purified by Percoll (Pharmacia AB) gradient centrifugation. Cell purity (> 90%) was verified by FACS staining with 5T2MM anti-idiotypic antibodies (Vanderkerken et al., 1997). 5T2MM cells were used for in vitro and in vivo experimental assays.

Peptides

LN, extracted and purified by immunoaffinity from the Engelbreth-Holm-Swarm (EHS) tumour was purchased from Sigma-Aldrich and peptide 11, a synthetic nonapeptide derived from the B1 chain of LN was purchased from Eurobiochem. A control scrambled nonapeptide (Ala-Asp-Cys-Gly-Gly-Ile-Pro-Ser-Tyr) was purchased from Genosys.

RNA extraction and Northern blotting

Northern blot hybridisation has been used to assess the expression of mRNA for the 37 kD precursor of 67LR from total RNA extracts from cultured HMCL. The plasmid DNA which carries the cDNA for the 37 kD precursor of 67LR (a kind gift from Dr Vincent Castronovo, Metastasis Research Laboratory, University of Liege, Belgium) was digested with XbaI. The released fragment was purified using the Wizard DNA clean up purification kit (Promega). Total RNA was isolated from cultured HMCL with the RNeasy Mini Kit (Qiagen). For Northern blot analysis, equal amounts of total RNA (10 μg) were denatured with 37% formaldehyde and denatured formamide, subsequently separated through a 1% agarose-formaldehyde gel and blotted onto Biobond nylon membranes (Sigma-Aldrich). cDNA probes encoding for the 37 kD precursor of 67LR were radioactively labelled with 32P-dCTP. Prehybridisation was performed in 50% formamide, 5 × Denhardt’s solution, 2 × SSC, 1% SDS and 100 mg ml⁻¹ denatured salmon sperm DNA for 3 hours at 42°C. Blots were subsequently hybridised overnight at 42°C. After hybridisation, membranes were stringently washed by vigorous rotation at 42°C for respectively 5, and 2 × 30 minutes in each of the following solutions: 2 × SSC/0.1% SDS and 0.2 × SSC/0.1% SDS. After washing, membranes were exposed to film (Kodak Biomax MR) overnight at ~80°C. As a control for the quality and quantity of the RNA samples analysed, the same blots were hybridised with a glyceraldehyde-3-phosphate dehydrogenase (GAPDH) cDNA probe. The blotting experiment was repeated twice with different RNA preparations.

Flow cytometric analyses

Fluorescence-activated cell sorting (FACS) analyses were performed to characterise the expression of 67LR on HMCL. After washing, cells were incubated with a monoclonal antibody (MLuC5) (IgM) recognising 67LR (kindly provided by Dr V Castronovo, Metastasis Research Laboratory, University of Liege, Belgium) and a polyclonal rabbit antiserum against 67LR (provided by Dr HK Kleiman, Institute of Dental and Craniofacial Research, National Institute of Health, Bethesda, USA). The last mentioned antiserum was also used to demonstrate 67LR expression on the murine 5T2MM cell line. As negative control, cells were incubated with isotype-matched irrelevant mouse Ig or rabbit Ig antibodies. Cells were subsequently washed in PBS/BSA and incubated with rabbit anti-mouse Ig G mouse serum or swine anti-rabbit IgG antiserum, both conjugated with fluorescein isothiocyanate (FITC) (Dako). Next, cells were washed, resuspended in PBS/BSA and finally analysed by flow cytometry (EPICS XL, Coulter Electronics). 67LR expression on MM plasma cells from patient samples was evaluated by a double staining procedure. Mononuclear cells isolated by ficoll centrifugation were incubated with the MLuC5 monoclonal antibody as indicated above. After washing, cells were simultaneously incubated with a PE conjugated goat antimouse IgM antiserum (Southern Biotechnology) and a Cy-5 conjugated CD38 specific antibody (HIT2) (BD Pharmingen). Plasma cell identification was controlled by co-expression of CD38 and intracytoplasmic κ/λ immunofluorescence using three-colour staining (rabbit anti-human IgG-FITC and Igλ-PE F(ab)₂ fragments (Dako)).
Co-culture of MM plasma cells with an endothelial cell line

In order to evaluate whether the expression of 67LR on MM plasma cells could be modulated by exposure to endothelial cells, mononuclear cells isolated from MM BM samples were co-cultured for 24 hours with the BM endothelial cell line 4LHBMEC (kindly provided by Dr A Dräger, Department of Hematology, Free University Amsterdam, The Netherlands). This cell line was cultured in 10% FCS medium 199 with 10 ng ml⁻¹ ECGF (Roche) until a confluent adherent monolayer was obtained. Mononuclear cells were incubated at 10⁶ cells ml⁻¹ in 5% FCS-RPMI with or without pre-established confluent monolayers of 4LHBMEC. After 24 hours non-adherent cells were removed by gentle pipetting and FACS analysis was performed.

In vitro cell migration assays

We performed in vitro migration assays with murine and human MM cells using an in vitro Transwell® cell migration system (Costar Corning) with 5 and 8 μm pore size membrane filters respectively. MM cells (1 × 10⁶) were added to the upper well in 100 μl medium. LN was diluted in 300 μl RPMI 1640 medium in varying concentrations and added to lower wells. Cells were tested for their spontaneous motility in response to control medium (RPMI 1640). The migration responses to LN were tested with controls in the same assay. After an incubation of 3 hours at 37°C in a humidified 5% CO₂ atmosphere, the number of cells that migrated through the filter into lower wells was quantified with a non-radioactive, colorimetric assay system using XTT (Roche) (Roehm et al, 1991). XTT with phenazine methosulfate (PMS) was added in each well after removal of inserts and incubated for 5 hours at 37°C. The absorbance of converted stain was measured spectrophotometrically with a 96-well microplate reader (Ceres 900 Bio-Tek International Inc) at a wavelength of 450 nm with a reference wavelength of 650 nm. Migration responses were determined as the percentage increase in cell migration compared to spontaneous migration. All experiments were performed in triplicate. Zigmund–Hirsh checkerboard analysis (Zigmund and Hirsh, 1973) was performed to distinguish chemokinetics from chemotaxis. For migration-inhibition experiments, MM cells were incubated with peptide 11 or with a blocking polyclonal antisera directed against 67LR (Clement et al, 1990). Scrambled peptide 11 and a polyclonal rabbit IgG (Chemicon International) served as controls. LN was used at 50 μg ml⁻¹. Migration responses were compared with control migration to LN and determined as described above.

For transendothelial migration, 2 × 10⁵ endothelial cells (4LHBMEC) were seeded on Transwell® filters. Confluent monolayers were formed after two days. Freshly isolated primary MM cells (1 × 10⁵) were added to the upper compartment of the Transwell® system, with or without peptide 11 (50 μg ml⁻¹), and allowed to migrate overnight. LN (50 μg ml⁻¹) was added as chemooattractant to the lower compartment. Transmigrated MM cells were recovered and counted by flow cytometry (FACSort, Becton Dickinson). Migration responses were determined as described.

In vivo experimental assays

For in vivo experiments, 5T2MM cells were labelled with 500 μCi ⁵¹Cr-chromate (Amersham). After washing, cell viability was assessed by trypan blue exclusion and cells were resuspended in RPMI 1640 at 2.5 × 10⁶ ml⁻¹. Peptide 11 or scrambled peptide 11 was solubilised at 2 mg ml⁻¹ in PBS and mixed with 5T2MM cells suspensions (5 × 10⁶) in a final volume of 200 μl. After 5 min incubation, 5T2MM cells were co-injected with peptide 11, scrambled peptide 11 or PBS intravenously into the tail vein of syngeneic mice. After 18 hours, mice were killed, legs, ribs and vertebrae were removed, subsequently rinsed in PBS and the radioactivity was determined by gamma-counting (MR48OC/TP ITM, Van Hopplynus) to estimate the level of tumour infiltration in the BM (Vanderkerken et al, 2000). Control and treatment groups each contained 9 mice.

Statistical analysis

Statistical analyses were performed using the 2-sample t-test. A P value of less than 0.05 was considered significant. Values were expressed as the mean ± SD.

RESULTS

LN receptor expression by HMCL, murine 5T2MM cells and fresh MM plasma cells

To characterise the HMCL with respect to their expression of 67LR, we performed Northern blotting and flow cytometric analysis using a specific monoclonal antibody to 67LR. Northern blot analysis revealed a homogenous expression of mRNA for the 37 kD precursor of 67LR in all 3 HMCL tested (Figure 1A). Flow cytometry demonstrated the surface expression of 67LR on HMCL and the murine 5T2MM cells (Figure 1B). 67LR expression on CD38⁺ plasma cells from 5 MM BM samples was more weak (Figure 1B). However, an increased expression of this receptor could be observed after direct exposure of tumour cells to a BM endothelial cell line (4LHBMEC) as illustrated in 3 patients (Figure 2).

Effect of LN on the in vitro migration of HMCL and murine 5T2MM cells

The migration of 3 HMCL (MM5.1, U266 and MMS.1) and the 5T2MM cells was evaluated in the presence of LN. As shown in Figure 3, increasing concentrations of LN (10 to 100 μg ml⁻¹) resulted in an increased number of cells migrating through the microporous membrane, with a mean increased migration response up to 90%. Checkerboard analyses revealed that MM cell migration to LN was chemotactic with a chemokinetic component (Table 1).

Inhibitory effect of peptide 11 on MM cell migration to LN

Peptide 11 has been shown to possess the ability to inhibit the biological activity of LN by competing for binding with 67LR. We investigated whether peptide 11 could modulate the interactions between MM cells and LN and we tested its effect on the migration of MM.5.1 and 5T2MM cells in response to LN. In vitro cell migration assays were carried out with LN (50 μg ml⁻¹) added to the lower well of the migration system. MM cells were placed in upper wells, together with peptide 11 (10, 50 or 100 μg ml⁻¹) or scrambled peptide 11 (100 μg ml⁻¹). As shown in Figure 4, peptide 11 inhibited MM cell migration to LN.
However, with increasing concentrations of peptide 11, inhibition of cell migration was gradually reduced. This last effect most likely relates to the chemokinetic effect of peptide 11 (Graf et al, 1987a). Scrambled peptide 11 did not affect in vitro MM cell migration to LN.

**Figure 1**  Expression of 67LR on HMCL, murine 5T2MM cells and MM cells from BM samples. (A) Representative Northern blot signals to transcripts from three HMCL show a homogeneous expression of the 37 kD precursor of 67LR in all tested HMCL. (B) Representative results of flow cytometric analysis are shown and expressed as log fluorescence versus number of cells. Background fluorescence is represented by white histograms, 67LR expression by black histograms. 67LR expression on malignant plasma cells from five bone marrow samples of MM patients was analysed by two-colour immunofluorescence using CD38-Cy5

**Effect of a blocking antiserum against 67LR on MM cell migration to LN**

In order to evaluate the importance of 67LR in the interaction of MM cells with LN, we tested the effect of a blocking polyclonal
antiserum against 67LR on MM cell migration to LN. As shown in Figure 5, antiserum against 67LR partially inhibited the migration response induced by LN in a dose-dependent manner.

Effect of LN and peptide 11 on the in vitro migration of primary MM cells

Previously, in vitro cell migration experiments to laminin were performed using primary MM cells freshly isolated from the bone marrow of patients with myeloma. Only minor migration responses were observed when LN was added to the lower compartment (data not shown). Considering the weak basal expression of 67LR on primary MM cells and its up-regulation when co-cultured with bone marrow endothelium, we carried out in vitro migration assays after coating the Transwell® filters with BM endothelial cells. In this condition, a significant migration response was observed to LN, indicating that 67LR was up-regulated by contact with endothelium, as found also in the coculture experiments. LN-induced migration of primary MM cells could also be inhibited by peptide 11, again suggesting a role of 67LR in the observed migration responses (Fig. 6).

Effect of peptide 11 on the in vivo migration of murine MM cells

The 5T2MM murine model was also used to evaluate whether treatment with peptide 11 affects the in vivo migration of tumour cells to the BM. Therefore, we injected $^{51}$Cr-labelled 5T2MM cells

Figure 2. Modulation of 67LR expression on MM plasma cells after exposure to the BM endothelial cell line 4LHBMEC. 67LR expression on malignant plasma cells was analysed before and after co-culturing mononuclear cells from three MM bone marrow samples with the 4LHBMEC cell line. Cells were analysed by two-colour immunofluorescence using CD38-Cy5.

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together with peptide 11 or scrambled peptide 11 intravenously into the tail vein of syngeneic mice. Tumour cell infiltration in the BM was determined by counting the radioactivity in legs, ribs and vertebrae after 18 hours. Results were represented as the % of recovered cpm in control and treated animals. As shown in Figure 7, co-injection of 5T2MM cells with peptide 11 significantly reduced (25%, P < 0.05) MM cell migration to the BM compartment. Scrambled peptide 11 did not affect the in vivo MM cell migration.

**DISCUSSION**

The homing of circulating MM cells to the BM implicates that these cells must be equipped with appropriate cell surface receptors, which allow binding to and traversing through the endothelium. Very little information is available about the molecular mechanisms used by MM cells during extravasation. Several studies indicated that the interaction of tumour cells with components of the ECM represents a key step in the biology of cancer cell migration and invasion. In the present study, we focused on the specific role of LN in MM cell migration. 3 well-characterised
Effect of LN on the in vitro transendothelial migration of primary
MM cells. For transendothelial cell migration, Transwell® filters were coated
with bone marrow endothelial cells as described in Materials and Methods.
LN was added into lower wells at 50 µg ml⁻¹. As negative control, dilution
medium (RPMI) was added to lower wells. Freshly isolated myeloma plasma
cells from 2 patients (pt 3 and pt 5) were placed into upper wells, with or
without peptide 11 (50 µg ml⁻¹). Columns represent the percentage increase
of control migration

Figure 6 Effect of LN on the in vitro transendothelial migration of primary
MM cells. For transendothelial cell migration, Transwell® filters were coated
with bone marrow endothelial cells as described in Materials and Methods.
LN was added into lower wells at 50 µg ml⁻¹. As negative control, dilution
medium (RPMI) was added to lower wells. Freshly isolated myeloma plasma
cells from 2 patients (pt 3 and pt 5) were placed into upper wells, with or
without peptide 11 (50 µg ml⁻¹). Columns represent the percentage increase
of control migration

Figure 7 Effect of peptide 11 on the in vivo migration of 5T2 MM cells.
5T2MM cells were labeled with ⁵¹Cr, mixed with peptide 11 (2 mg ml⁻¹) or
scrambled peptide 11 and injected intravenously into the tail vein of
syngeneic mice. After 18 hours, mice were killed; legs, ribs and vertebrae
were removed, subsequently rinsed in PBS and radioactivity was determined
by gamma counting. Control and treatment groups contained 8 mice. Each
column represents the mean percentage ± SD of total recovered radioactivity
in the bone marrow. The P value was calculated from the 2- sample
t-test (*: statistically significant)

HMCL, as well as the murine 5T2MM cell line were selected for
our experiments. The expression of 67LR, a high-affinity LN-
binding protein, was analysed by Northern blotting and FACS
analysis. All human MM cell lines and the murine 5T2MM cells
expressed 67LR. 67LR is expressed by several normal cell types,
as well as cancer cells of different origin, including B cell
lymphoma (Carbone et al, 1995), but its expression on MM cells
has not been reported so far. In vitro migration experiments
demonstrated that LN stimulated the chemotactic migration of
HMCL and murine 5T2MM cells. Previous studies revealed that
the action of LN appeared through binding to 67LR with a
sequence of 9 amino acids Cys-Asp-Pro-Gly-Tyr-Ile-Gly-Ser-Arg
(CDPGYIGSR) (Situ et al, 1984). A synthetic nonapeptide
(peptide 11), derived from this aminosequence was found to be
able to compete with LN for binding to 67LR (Graf et al, 1987b).
In vivo, it reduces the formation of experimental lung and oste-
olytic metastasis (Iwamoto et al, 1987, 1996; Nakai et al, 1992)
and inhibits human pre-B leukaemic cell growth and dissemination
to organs in SCID (Yoshida et al, 1999). We evaluated the effect of
this synthetic nonapeptide on MM cell migration and observed that
peptide 11 reduced MM cell migration to LN. In addition, we
could demonstrate that the migration of MM cells could be inhib-
ited by a blocking polyclonal antiserum against 67LR (Clement
et al, 1990). These data allow us to conclude that 67LR plays an
important role in the motility-promoting activity of LN on MM
cells. The polyclonal antiserum against 67LR could not induce a
complete inhibition of MM cell migration to LN indicating that
other receptors might influence this motility process as well. In a
previous study, Shibayama et al (1995) demonstrated that the
migration of 2 HMCL (FR4ds and OPM-1 ds) to LN could be
partially inhibited by a blocking antibody against the common
chain of β1 integrins. This may indicate that the overall migration
of MM to LN might be mediated by multiple receptors.

Interestingly, 67LR expression could be enhanced when primary
MM cells were co-cultured with a human BM endothelial cell line,
whereas 67LR expression remained unchanged after contact with
BM stromal fibroblasts (data not shown). Moreover, in vitro
migration experiments with primary MM cells revealed an
enhanced migration response to laminin when Transwell® filters of
the migration system were coated with endothelial cells. This
could indicate that 67LR is temporally up-regulated at the moment
that MM cells adhere to and migrate through the endothelium
while receptor expression decreases when tumour cells reside as
fully matured plasma cells in the BM compartment. Accordingly,
HMCL might express more 67LR because they reflect a more
immature tumour stage than fully matured plasma cells. It was
demonstrated that 67LR was co-localised with alpha-actin and
vinculin, 2 structural cell proteins (Massia et al, 1993) suggesting a
direct role for 67LR in cell motility.

To determine at which level LN-67LR interactions also influ-
ence the in vivo homing of MM cells, we used the murine 5T2MM
model. This model has been intensively characterised and was
found to express very similar pathological and biological features
as human MM (Vanderkerken et al, 1996, 1997). In a recent study,
we could demonstrate that 5T2MM cells in vivo selectively home
to BM and liver and not to other organs. Moreover, we found
that 5T2MM cells adhere selectively to BM, spleen and liver and not to
lung endothelial cells (Vanderkerken et al, 2000). In the present
study, we demonstrated that 5T2MM cells express 67LR and
migrate in vitro to LN and peptide 11. When radiolabelled
5T2MM cells were mixed with peptide 11 and co-injected into
syngeneic mice, the proportion of tumour cells which accumulated
in the BM after 18 hours was significantly decreased in peptide 11-
treated animals. This observation indicates that interaction
between 67LR and LN is at least partially involved in the in vivo
migration of 5T2MM cells through BM endothelium. Because of
the broad expression of LN in extracellular matrices throughout the body, it is clear that this molecule itself can not be the only factor that determines the specificity of MM cell homing. It can be assumed that several adhesion mechanisms and chemotactic signals co-act to enhance the selectivity of tumour cell migration. Moreover, the restricted localisation of MM cells in the BM might also relate to the presence of a unique combination of growth and survival factors, which is not available at other tissue sites. In conclusion, we demonstrated that MM cells express the high-affinity 67 kD LN receptor and that this receptor is involved in LN-mediated cell migration. It is likely that 67LR is involved during the extravasation of MM cells to the bone marrow. The homing of myeloma cells to the bone marrow is believed to be a multistep process, involving the action of various adhesion molecules, chemotactic signals and metalloproteases. Interference with one of the key steps in this process (like migration to laminin) might be an important therapeutic tool to reduce tumour dissemination.

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