Single-Fluorescent Protein Reporters Allow Parallel Quantification of Natural Killer Cell-Mediated Granzyme and Caspase Activities in Single Target Cells

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Natural killer (NK) cells eliminate infected and tumorigenic cells through delivery of granzymes via perforin pores or by activation of caspases via death receptors. In order to understand how NK cells combine different cell death mechanisms, it is important to quantify target cell responses on a single cell level. However, currently existing reporters do not allow the measurement of several protease activities inside the same cell. Here, we present a strategy for the comparison of two different proteases at a time inside individual target cells upon engagement by NK cells. We developed single-fluorescent protein reporters containing the RIEAD or the VGPD cleavage site for the measurement of granzyme B activity. We show that these two granzyme B reporters can be applied in combination with caspase-8 or caspase-3 reporters. While we did not find that caspase-8 was activated by granzyme B, our method revealed that caspase-3 activity follows granzyme B activity with a delay of about 6 min. Finally, we illustrate the comparison of several different reporters for granzyme A, M, K, and H. The approach presented here is a valuable means for the investigation of the temporal evolution of cell death mediated by cytotoxic lymphocytes.

Keywords: natural killer cells, cytotoxic lymphocytes, single-fluorescent protein reporters, granzyme and caspase activity, apoptosis and cell death

INTRODUCTION

As part of the innate immune system, natural killer (NK) cells can eliminate infected and tumorigenic cells (1). To do so, they adhere to a target cell and establish an immunological synapse (2). The following NK cell receptor signaling can trigger the release of cytotoxic granules from the NK cell into the cleft of this synapse (3). Like cytotoxic T lymphocytes (CTLs), NK cells have two mechanisms to induce cell death of target cells. In the first mechanism, granzymes are released from cytotoxic granules and enter the target cell via perforin pores and induce cell death. In the second mechanism, CD95L or TRAIL are presented at the surface of NK cells and induce extrinsic apoptosis in target cells through activation of the death receptors CD95 or TRAIL-R1/-R2 (4, 5).
How NK cells orchestrate the activities of granzymes and the activation of extrinsic apoptosis remains poorly understood. Extrinsic apoptosis starts with the formation of the so-called death-inducing signaling complex, composed of activated death receptors and recruited FADD adaptor proteins and initiator procaspases-8/-10. Once activated, these caspases cleave and activate effector procaspase-3/-7 (6, 7), leading to apoptosis, unless presence of XIAP blocks their activity (8, 9). When the pro-apoptotic Bcl-2 protein BID is cleaved by caspase-8/-10 in sufficient amount, truncated BID induces mitochondrial outer membrane permeabilization. Subsequent release of cytochrome c activates caspase-9, while release of SMAC induces the degradation of XIAP, both leading to massive activation of effector caspases.

To deliver granzymes in the cytosol of target cells, perforin forms a pore in cellular membranes (10). It is debated if this occurs at the plasma membrane (11, 12) or the membrane of endosomes (13–15). Of the five human granzymes A, B, H, K, and M, granzyme B is the best characterized one and shares substrate specificity with caspases for cleavage after aspartate residues (16–18). Both, granzyme B and caspase-8 can cleave BID, yet, at different sites, at D75 (RIEAD’S) and D60 (ELQTD’G) (19), respectively. While granzyme B has been shown to cleave the initiator procaspase-8 (20) and the effector procaspase-3 (21–25), other substrates measured in vitro have been reported to be more efficiently cleaved, for example, DNA-PKc or BID (23, 26–28). From this perspective, granzyme B is suggested to play a role not only as an initiator but also as executioner enzyme in target cell death (9).

Having reporters that would allow the measurement of the contribution of granzymes and caspases in a single cell would be beneficial to characterize the activity of NK cells. Specific protease biosensors based on luciferase (29, 30), fluorophore quenching (31), and FRET (32, 33) (Table 1) have facilitated the study of the killing mechanism by granzymes and death receptors. However, they do not easily allow multiplexing for the quantification of several protease activities in single cells. Parallel assessment of protease activity inside single cells would allow for a better understanding of the temporal order of signaling events in the NK cell killing mechanism. In order to reach this aim, we present an approach to measure NK cell-mediated activity of two proteases at once in single target cells. We demonstrate our approach by measuring granzyme B, caspase-8, and caspase-3 activity in target cells exposed to NK cells. The pallet of reporters can easily be extended by cloning cleavage linkers, as illustrated here with the measurement of potential substrates for different granzymes. We believe that these reporters offer a valuable resource to characterize the physiology of NK cells or to test the activity of patient-derived NK cells.

### RESULTS

#### Measurement of Protease Activity Induced by NK Cells in Living Target Cells

To get insights into the process by which individual NK cells kill their target cells, we aimed at comparing the strength of different proteases inside single target cells. To achieve this, we designed fluorescent reporters following the strategy that we previously established to measure caspase activity upon CD95 activation (34). These reporters consist of one fluorescent protein fused to a localization domain through a linker that can be specifically cleaved by proteases. In this study, we use the nuclear export signal (NES) as localization domain (Figure 1). As exemplified below, this reporter allows an easy quantification of protease activity, and it detects any activity originating from an enzyme facing the cytosol. Protease cleavage leads to separation of the fluorescent protein from the localization domain. The free fluorescent protein is small enough to enter or exit the nucleus by passive diffusion, a process that takes about 1 min. Reporter cleavage can be quantified by measuring the increase of fluorescence signal in the nucleus. This is ideally imaged by time-lapse fluorescence confocal microscopy as this allows the measurement of the fluorescence intensity inside the nucleus without contamination of signal from the cytosol below and above. Thus, by quantifying the spatial redistribution of the fluorescence signal inside the target cell, reporter cleavage can be calculated. Image analysis can be done using the freely available ImageJ software and proceeds as follows: generation of an image stack from time series data, background subtraction, and measurement of the mean fluorescence signal in a region of interest representing the cell nucleus. To estimate the extent of substrate cleavage, this nuclear signal is normalized to the

### Table 1

| Substrate/ cleavage site | Cleaved by protease | Protein containing the cleavage site | Reference |
|--------------------------|---------------------|-------------------------------------|-----------|
| ELQTD’G                  | Caspase-8/-10       | Human BID                           | (19, 34)  |
| RIEAD’S                  | Granzyme B,         | Human BID                           | (19, 29, 35–37) |
|                          | Caspase-10          |                                     |           |
| DEVD’R                   | Caspase-3/-7        | PARP-1                              | (29, 38)  |
| VGPD’FGR                 | Granzyme B          | DNA-PK                              | (29, 31–33, 35) |
| RIEPO’S                  | Granzyme B          | Mouse BID                           | (26, 39)  |
| IGNR’S                   | Granzyme A          |                                     | (39)      |
| PTSY’G                   | Granzyme H          |                                     | (39)      |
| YRFK’G                   | Granzyme K          |                                     | (39, 40)  |
| KVPL’AA                  | Granzyme M          |                                     | (39, 41)  |
| DVAHK’QGL                | Granzyme A          | NDUFS3                              | (42)      |

FIGURE 1 | Schematic representation of a single-fluorescent protein (single FP) reporter and its fluorescence distribution inside the cell when using a nuclear export signal as localization domain. Upon cleavage of the reporter, the fluorescent protein can diffuse passively into the nucleus.
cytoplasmic one. One possibility is to perform this normalization for each time point, which requires more work but has the advantage of correcting for potential photobleaching over time:

\[ I_{\text{normalized}}(t) = \frac{I_{\text{nuclear}}(t)}{I_{\text{cytoplasm}}(t)} \]

In this case, the normalized intensity tends toward 1, or 100%, when cleavage is complete. The data presented here were analyzed in this way. The second possibility consists of normalizing with the cytosolic signal before the addition of the NK cells. This approach is relevant if photobleaching can be neglected:

\[ I_{\text{normalized}}(t) = \frac{I_{\text{nuclear}}(t)}{I_{\text{cytoplasm}}(t = 0)} \]

In this case, as the nucleus and the cytoplasm are roughly occupying the same volume, the normalized intensity tends toward 0.5 or 50%.

Since each reporter contains only one fluorescent protein, these so-called single-fluorescent protein reporters allow parallel assessment of several reporters within the same cell. Hence, different comparison can be realized: the measurement of (i) different cleavage sites for different proteases or of (ii) different cleavage sites for the same protease.

**Granzyme B Activity Can Be Measured With Single-Fluorescent Protein Reporters Having the RIEADS or the VGPD Cleavage Site**

In order to establish single-fluorescent protein reporters for the measurement of granzyme B activity, we designed and tested two different reporters carrying a linker sequence known to be cleaved by granzyme B. The first reporter carries the amino acid sequence RIEADS (single amino acid code), which is present in the protein BID. The second reporter carries the amino acid sequence VGPD from the protein DNA-PKc as cleavable linker. We co-expressed the two reporters NES-RIEADS-mCherry and NES-VGPD-mGFP in HeLa cells expressing CD48, a ligand for the activating NK cell receptor 2B4 (CD244), which renders them more sensitive to killing by NK cells (43). The transfected target cells were imaged by confocal laser scanning microscopy over time upon addition of the human NK cell line NK92-C1. Cell death was recognized from images by cell rounding and cell shrinkage following NK cell engagement (Figure 2A). About 20 min before target cell death, NES-RIEADS-mCherry and NES-VGPD-mGFP reporter cleavage was detected from the appearance of fluorescence inside the nucleus (Figure 2A). In order to plot the cleavage kinetics of several cells independently of the time of NK cell engagement, we defined the time of death and caspase-8 activity in a control experiment using IZsCD95L as inducer (Figure 4A). The nuclear protein H2B-eBFP2 served as a fluorescent marker to recognize cells that contain the plasmid encoding shRNA (Figure 4B). Upon addition of NK cells to HeLa (CD48) cells expressing control shRNA, we observed NES-RIEADS-mCherry reporter cleavage as expected indicating granzyme B activity. In addition, the NES-ELQTD-mGFP reporter was cleaved only up to 10% (Figure 3A) and the NES-VGPD-mCherry reporter was virtually not cleaved (Figure 3B). This shows that granzyme B activity (using either the RIEADS- or the VGPD-reporter) can be clearly distinguished from caspase-8 activity (using the ELQTD-reporter) within the same cell.

**Single-Fluorescent Protein Reporters Allow Multiplexing: Distinguishing Different Proteases Inside the Same Cell**

CD95 signaling in HeLa cells leads to notable caspase-8 and -3 activation (34). On the one hand, it was reported that caspase-8 can get activated through cleavage by granzyme B (20). On the other hand, it was shown that granzyme B activates effector caspases through cleavage of BID (24, 25, 27, 28). Deciphering the contribution of caspase-8 and granzyme B to NK cell-mediated cell death, and in particular, the potential activation of caspase-8 by granzyme B, is an intriguing question that could help to better understand cell death signaling by NK cells.

Therefore, we tested if caspase-8 and granzyme B activity can be distinguished within the same cell using our single-fluorescent protein reporters. For this aim, we measured NES-ELQTD-mGFP (for caspase-8), together with either NES-RIEADS-mCherry or NES-VGPD-mCherry (for granzyme B). Upon addition of soluble trimerized CD95L (IZsCD95L) to HeLa cells, we observed efficient cleavage of the caspase-8 reporter NES-ELQTD-mGFP. In contrast, in the same cells, the NES-RIEADS-mCherry reporter was cleaved only up to 10% (Figure 3A) and the NES-VGPD-mCherry reporter was virtually not cleaved (Figure 3B). This shows that granzyme B activity (using either the RIEADS- or the VGPD-reporter) can be clearly distinguished from caspase-8 activity (using the ELQTD-reporter) within the same cell.

Transient expression of shRNA against CD95 led to a reduction of CD95 protein expression on the surface of HeLa cells (Figure S1 in Supplementary Material) and furthermore to absence of cell death and caspase-8 activity in a control experiment using IZsCD95L as inducer (Figure 4A). The nuclear protein H2B-eBFP2 served as a fluorescent marker to recognize cells that contain the plasmid encoding shRNA (Figure 4B). Upon addition of NK cells to HeLa (CD48) cells expressing control shRNA, we observed NES-RIEADS-mCherry reporter cleavage as expected indicating granzyme B activity. In addition, the NES-ELQTD-mGFP reporter was cleaved on average up to 7% indicating an additional activation of caspase-8 (Figure 4C). This caspase-8 activity could be due to the activation of caspase-8 by granzyme B, or due to the activation of death receptors during the engagement by the NK cells. To distinguish these two possibilities, we repeated the experiment in HeLa cells expressing a shRNA against CD95. The absence of CD95 expression abolished NES-ELQTD-mGFP reporter cleavage (Figure 4D). This demonstrates that the caspase-8 activity was due to activation of the CD95 pathway by NK cells, and it also reveals that caspase-8 is not, or not efficiently, activated by granzyme B. When using activated primary human NK cells as effectors,
we saw a similar cleavage of the NES-VGPD-mCherry granzyme B reporter and of the NES-ELQTD-mGFP caspase-8 reporter in HeLa cells and in MDA-MB-468 breast carcinoma cells (Figure S1 in Supplementary Material). This demonstrates that NK cells use granzyme B dependent (via cytotoxic granules) and caspase-8-dependent (via CD95) pathways to kill target cells.
It is known that effector caspase-3 activation can be a consequence of granzyme B activity (26). We thus hypothesized that caspase-3 activity should occur with a delay after granzyme B activity, as in the case of CD95-mediated extrinsic apoptosis (7, 34). To test this, we measured reporter cleavage of NES-VGPD-mCherry for granzyme B and NES-DEVDR-GFP for caspase-3 in HeLa cells upon addition of NK cells (Figure 5A). We found that caspase-3 activity appears on average about 6 min later than granzyme B (Figure 5B). The onset of caspase-3 activity was about 10 min before cell death, highlighting the speed of this cellular process. These data also show how precisely cell death can be analyzed on the single cell level with this approach using single-fluorescent protein reporters.

Testing Potential Reporters for Different Granzymes

Natural killer cells notably express granzyme A and B (45, 46), but they also express the less characterized granzymes H, K, and M (47). In order to further illustrate our approach and to potentially capture the activity of granzymes A, M, H, and K in single target cells, we designed fluorescent reporters that contain cleavage site candidates based on existing data from literature (Table 1). Using positional scanning combinatorial libraries, optimal substrates for the five human granzymes have been previously identified and characterized (39). Among them, the IEPD and IGNR sequence have been further developed as fluorescent label and inhibitor for granzyme B and granzyme A (39). Moreover, optimal substrates were determined to be YRFK for granzyme K, KVPL for granzyme M, and PTSY for granzyme H (39). In this study, we additionally designed a reporter containing the sequence DVAHKQL, which is present in the mitochondrial complex I protein NDUFS3, since granzyme A was reported to cleave this protein at this site after the amino acid lysine in this sequence (42). We measured these six different reporters in comparison to the granzyme B reporters NES-VGPD-mCherry or NES-RIEADS-mGFP. We found that the RIEADS and VGPD reporters were more sensitive to detect granzyme B activity compared to the IEDP reporter (Figure 6A). Also, granzyme B activity was clearly dominant among the different reporters. Moreover, no cleavage of DV AHQL (granzyme A) and YRFK (granzyme K) reporter cleavage was observed. This comparison of different reporters would suggest that granzyme B and granzyme A are the most abundant granzymes used by the NK92-C1 cell lines to cleave substrates in the cytosol of target cells. This experiment shows how single-fluorescent protein reporter can be used to examine protease activity that is either induced inside target cells or stemming from cytotoxic lymphocytes such as NK cells.

**DISCUSSION**

The single-fluorescent protein reporters employed in this study are part of a larger panel of protease reporters, based, for
FIGURE 5 | Measurement of granzyme B and caspase-3 activity in single target cells. (A,B) HeLa cells transfected with CD48, NES-VGPD-mCherry (for granzyme B), and NES-DEVDR-mGFP (for caspase-3) were imaged by confocal microscopy upon incubation with NK92-C1 cells (E:T = 2). (A) Example images of the time series. Note that the fluorescence signal inside the nucleus is visible for NES-VGPD-mCherry at t = 50 min and for NES-DEVDR-mGFP at t = 56 min. Scale bar: 10 µm. (B) Quantification of reporter cleavage of 23 cells and boxplot showing the time of target cell death. On average, caspase-3 activity appears with a short delay of about 6 min after granzyme B activity.

example, on cyclic luciferase (29, 48), fluorophore quenching (31), FRET imaging (32, 33, 49), or subcellular localization of a fluorophore (34, 50). Our reporters provide us with the possibility to precisely investigate NK cell-induced cell death since they allow the measurement of two protease activities within the same cell. This feature is useful to correlate the
activity of different enzymes within single cells and, therefore, independently of the large cell-to-cell variability due to the stochasticity of the NK–target cell contact. We used a confocal laser scanning microscope with a motorized stage that can record several user-defined fields of view one after the other. Thanks to that, we typically achieved the acquisition of around 40–50 single cells per experiment with a time-resolution of 2–4 min. The experimenter should find an optimal compromise between the number of imaged cells and the time resolution, which in turn depends on the speed of the autofocus and microscope type. In this context, a spinning disk microscope could provide higher imaging rate over a confocal laser scanning microscope.
Higher cell numbers per field of view may be also achieved by working with a target cell line that stably expresses the reporters. Regarding image analysis, we chose a simple manual workflow that we describe in detail in the Section "Materials and Methods," but we could also envision the development of automated image analysis workflows.

We demonstrate our approach using HeLa cells as targets as they are known to be responsive to death receptor ligands (34, 51) and granzymes (20, 27). Furthermore, adherent target cells such as HeLa cells are convenient for reporter expression and live microscopy. In particular, time-lapse microscopy of HeLa cells facilitates the identification of cell death, which was clearly visible from morphological changes like cell rounding, blebbing, and shrinkage. A limitation of the approach may be the use of target cells growing in suspension, as morphological changes due to cell death are not easily visible. We cannot exclude that one target cell was engaged by two or more NK cells at the same time, which may impact the kinetics of death signaling. Here, we worked with NK cell to target cell ratios in the range of 1/1 to 3/1. To ensure responses from single NK cells, we suggest as one possibility to add fewer NK cells, which should minimize the chance that several NK cells hit the same target cell.

We showed that the reporters developed here containing the RIEADS or the VGPD cleavage site are suitable for the measurement of granzyme B activity. We furthermore systematically tested putative reporters containing amino acid sequences that were previously shown to be cleaved by granzyme A, B, M, H, and K (39). Primary resting and activated NK cells were shown to express all granzymes, with the abundance order A, B > H > K, M (52). While granzyme B activity was the most prominent in our experiments, we could detect some activity of granzymes A, H, and K, but no activity of granzyme M. The reporters used here contain a nuclear export signal for the measurement of any enzyme activity in the cytosol. This may explain why the DVAHKQL reporter for granzyme A was not cleaved in target cells upon NK cell addition, since this substrate would naturally reside inside mitochondria (42). However, the contribution of other granzymes to cell death still remains to be further investigated, for example, by testing NK cells or CTLs at different maturation stages (53) or by taking into account the expression levels over long-term experiments (54).

Our experiments have demonstrated that granzyme B reporters can be applied in combination with caspase-8 or caspase-3 activity reporters. Strong caspase-3 activity correlated with granzyme B activity. These results support earlier studies (21, 23, 27, 28) showing that granzyme B can directly or indirectly (through BID) induce caspase-3 activation. Our data show that this is a fast process, with caspase-3 activity being detectable within 6 min after granzyme B activity. In contrast, caspase-8 activity was only detectable in target cells expressing the CD95 receptor but not in cells where CD95 was knocked-down, despite granzyme B activity. This suggests that there is no major activation of caspase-8 by granzyme B, in contrast to earlier studies (20). However, this also demonstrates that NK92 cells can indeed use two pathways to kill target cells: exocytosis of granzymes and perforin, resulting in detectable granzyme B activity inside the cytosol of target cells, and surface expression of CD95L and TRAIL, resulting in caspase-8 activity. Moreover, we also found clear reporter cleavage showing granzyme B and caspase-8 activity in HeLa and MDA-MB-468 breast carcinoma cells with activated primary NK cells as effector cells. This demonstrates the feasibility of the method for the study of different target cells and primary NK cells. Further work will be required to precisely understand how granzyme B and death receptor signaling are used by NK cells. We can envisage our reporters for the use in other target cells, where amounts of activating and inhibiting NK cell receptors are different or alternatively, where the amounts of death receptors or serpinB9 are different. Single-fluorescent protein reporters present a valuable tool to decipher cell death mechanisms induced by NK cells. We believe that this approach opens the door for the characterization of death receptor- versus granzyme-mediated target cell killing, in particular, the temporal evolution of these two death mechanisms in the context of serial killing by NK cells.

MATERIALS AND METHODS

Constructs

Human CD48 (UniProtKB P09326), human serpin B9/PI-9 (from Gateway cDNA library of the DKFZ), and IZ-sCD95L (55) were subcloned in the pIRES-puro2 vector. Fluorescent reporters were cloned based on constructs described previously (34) in pEGFP N1 and C1 vectors (Takara Bio Europe Clontech, Saint-Germain-en-Laye, France). DNA oligonucleotides encoding cleavage sites were cloned with AgeI/NotI or BsrGI/NotI (Table 2). The amino acid sequence starting from the N-terminus of the nuclear export signal (NES) is MNLVLDQKLEELEDEQQ. shRNA against CD95 and scrambled shRNA (see Table 2 for DNA sequences) were cloned in the pSilencer3.1 H1 Neo vector with HindIII and BamHI.

Cell Culture

HeLa and MDA-MB-468 cell lines were maintained in Dulbecco’s modified eagle medium (Invitrogen, Darmstadt, Germany), containing 10% fetal calf serum (Biochrom AG, Berlin, Germany), penicillin/streptomycin, 100 µg/ml each (Invitrogen) and 1% non-essential amino acids plus 1 mM Sodium Pyruvate (Gibco) for MDA-MB-468 only. HeLa (CD48) cells over-express CD48 and were maintained in medium supplemented with 0.5 µg/ml puromycin (Sigma-Aldrich). NK92-C1 that stably express IL-2 were maintained in phenol red-free Minimum Essential Medium Eagle Alpha (MEMx, Sigma-Aldrich) without ribonucleosides and deoxyribonucleosides but containing sodium bicarbonate, supplemented with 2 mM L-glutamine, 50 µM 2-mercaptoethanol, 12.5% horse serum, 12.5% fetal bovine serum, and penicillin/streptomycin, 100 µg/ml each (Invitrogen). Human NK cells were isolated from PBMCs using the Dynabeads Untouched Human NK Cell kit (Thermo Fisher Scientific) according to the manufacturer's instructions. NK cells were between 90 and 99% CD3−, CD56+, and NKp46+ as assessed by flow cytometry. NK cells were expanded in 96-well round-bottom plates (Nunc) with irradiated K562-mbIL15-41BBL feeder cells (kind gift from Dario Campana) in IMDM Glutamax supplemented with 10% FCS and 1% penicillin, IL-2 (100 U/ml, NIH Cytokine Repository),
ing secreted ligand was harvested. At the next day, supernatant containing IZsCD95L, cloned in pIRES-puro2, was produced in 293T cells, which were seeded in 6-well plates. 24 h after cell culture medium is found. From that position, an offset of a few micrometers was chosen to image a representative cross-section of the cell. We set the offset position so that an image of the cell is made approximately at the mid-height of the nucleus where the signal from the cytosol above and below the nucleus is minimal, and mostly cytosolic signal next to the nucleus is present. One image plane per field of view was acquired and cells were imaged once before the addition of NK cells. The resolution was 512 × 512 pixel and images were acquired in 8-bit or 16-bit. Fluorescence of mGFP and mCherry was acquired in line sequential mode. mCherry, we used the helium–neon laser (561 nm), detection range: 600–660 nm, for mGFP, we used the argon laser (488 nm), detection range: 500–560 nm. Cells were grown and imaged in 8-well ibidi chambers (ibidi GmbH, Planegg/Martinsried, Germany). We applied one-, two-, or threefold more effector NK cells compared to target cells (E:T = 1, E:T = 2, or E:T = 3, respectively), hence about 6 × 10⁴ to 1.8 × 10⁵ NK cells per well. One field of view in confocal microscopy contained about 2 to 6 transfected cells. All cells expressing both reporters were analyzed. The time-resolution was about 2–4 min depending on the number of imaged fields and microscopy settings. Equal target cell preparation, equal NK cell preparation (cell number counting, transfection), and parallel measurement of different wells of an 8-well chamber allowed comparison of different conditions in one experiment. Time-series data were acquired for up to 12 h.

**Image Analysis**

Images were analyzed using ImageJ (56). To quantify the nuclear redistribution of fluorescence intensity over time in single cells, the nuclear intensity was measured. For this, images of each channel were background subtracted. To subtract the background, we selected a region of interest where no cell was present and applied the plugin for background subtraction, which we provided online https://github.com/jbeaudouin/NK_cell. Alternatively, we suggest using the background subtraction function in the “Process” tab provided in ImageJ or Fiji, which is based on the rolling ball algorithm. In most cases, two reporters were measured within one cell: to analyze them, one channel was assigned green, the other red, both were transformed into RGB, and they were then superimposed. The mean intensity within the nucleus was quantified by choosing a representative region within the nucleus at each time point until cell death was observed. We chose to normalize the nuclear intensity to the cytosolic intensity at each time point for soluble-cytosolic reporters to calculate the percentage of reporter cleavage.

The detailed image analysis workflow is as follows:

1. Open image stack of the first channel (time-series of one field of view) in ImageJ or Fiji. If applicable, open image stack of the second channel (time-series of one field of view) in

| Cleavage sites: sense | Oligonucleotide, 5′ → 3′ |
|-----------------------|-------------------------|
| KVPLAA (as)           | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| KVPLAA (as)           | CGGCGGCTTCCTCAGGCGGGAAGCTGACACAATTTGCAAG |
| IEPDG (as)            | CCGGTGGGCGGATTGAAACAGATGTCGTGGAGGGG |
| IEPDG (as)            | CGGCGGCTCTCTCACCACCATGGGTAACAGTACGG |
| IGNRS (s)             | CCGGTGGGCGGATTGAAACAGATGTCGTGGAGGGG |
| IGNRS (s)             | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| PTSYG (s)             | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| PTSYG (s)             | CGGCGGCTCTCACCACCATGGGTAACAGTACGG |
| YRFKG (as)            | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| YRFKG (as)            | CGGCGGCTCTCACCACCATGGGTAACAGTACGG |
| DVAHKQL (s)           | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| DVAHKQL (s)           | CGGCGGCTCTCACCACCATGGGTAACAGTACGG |
| VGPDFGRG (s)          | GTACAAGGCGTAAGGCTGCTCCGGCGGAGGG |
| VGPDFGRG (s)          | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| RIEADS (s)            | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| RIEADS (s)            | CGGCGGCTCTCAGGCGGGAAGCTGACACAATTTGCAAG |
| shRNA (CD95)          | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| shRNA (scrambled)     | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |

and IL-15 (5 ng/ml, PAN-Biotech) at 37°C in a humidified 5% CO₂ incubator. IL-21 (100 ng/ml, Miltenyi Biotec) was added on the first day. Soluble CD95 ligand fused to the isoleucine-zipper domain (IZsCD95L, cloned in pIRES-puro2) was produced in 293T cells, which were seeded in 6-well plates. 24 h after cell transfection using JetPrime reagent (Polyplus), supernatant was replaced by fresh medium. At the next day, supernatant containing secreted ligand was harvested.

**Microscopy**

The presented approach uses fluorescence microscopy, which allows the extraction and correlation of several features, including the time of cell death. Time-lapse microscopy was performed with the TCS SP5 confocal laser scanning microscope from Leica (Leica Microsystems CMS GmbH, Mannheim, Germany) equipped with a 63×/1.4 OIL, HCX PL APO CS objective. We used the live data mode of the Leica software for autofocusing as described in Ref. (34). With the help of the autofocus option, the height position of the interface between the glass and the cell culture medium is found. From that position, an offset of a few micrometers was chosen to image a representative cross-section of the cell. We set the offset position so that an image of the cell is made approximately at the mid-height of the nucleus where the signal from the cytosol above and below the nucleus is minimal, and mostly cytosolic signal next to the nucleus is present. One image plane per field of view was acquired and cells were imaged once before the addition of NK cells. The resolution was 512 × 512 pixel and images were acquired in 8-bit or 16-bit. Fluorescence of mGFP and mCherry was acquired in line sequential mode. mCherry, we used the helium–neon laser (561 nm), detection range: 600–660 nm, for mGFP, we used the argon laser (488 nm), detection range: 500–560 nm. Cells were grown and imaged in 8-well ibidi chambers (ibidi GmbH, Planegg/Martinsried, Germany). We applied one-, two-, or threefold more effector NK cells compared to target cells (E:T = 1, E:T = 2, or E:T = 3, respectively), hence about 6 × 10⁴ to 1.8 × 10⁵ NK cells per well. One field of view in confocal microscopy contained about 2 to 6 transfected cells. All cells expressing both reporters were analyzed. The time-resolution was about 2–4 min depending on the number of imaged fields and microscopy settings. Equal target cell preparation, equal NK cell preparation (cell number counting, transfection), and parallel measurement of different wells of an 8-well chamber allowed comparison of different conditions in one experiment. Time-series data were acquired for up to 12 h.

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| DNA sequence encoding protease cleavage sites used for plasmid cloning. |
|--------------------------|--------------------------|
| KVPLAA (s)               | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| KVPLAA (as)              | CGGCGGCTTCCTCAGGCGGGAAGCTGACACAATTTGCAAG |
| IEPDG (as)               | CCGGTGGGCGGATTGAAACAGATGTCGTGGAGGGG |
| IEPDG (as)               | CGGCGGCTCTCTCACCACCATGGGTAACAGTACGG |
| IGNRS (s)                | CCGGTGGGCGGATTGAAACAGATGTCGTGGAGGGG |
| IGNRS (s)                | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| PTSYG (s)                | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| PTSYG (s)                | CGGCGGCTCTCACCACCATGGGTAACAGTACGG |
| YRFKG (as)               | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| YRFKG (as)               | CGGCGGCTCTCACCACCATGGGTAACAGTACGG |
| DVAHKQL (s)              | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| DVAHKQL (s)              | CGGCGGCTCTCACCACCATGGGTAACAGTACGG |
| VGPDFGRG (s)             | GTACAAGGCGTAAGGCTGCTCCGGCGGAGGG |
| VGPDFGRG (s)             | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| RIEADS (s)               | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| RIEADS (s)               | CGGCGGCTCTCAGGCGGGAAGCTGACACAATTTGCAAG |
| shRNA (CD95)             | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
| shRNA (scrambled)        | GATCCAATCTCATTCGATGCATACTTCAAGAGAGTGGAGGGG |
Assessment of the Time of Target Cell Death

Cell death of HeLa target cells could be determined by the change of the cell’s morphology. While cell rounding was often observed, we found also cells showing shrinkage without typical apoptotic blebbing. Changes in target cell morphology were unambiguously detected by monitoring living target cells by time-lapse microscopy in the fluorescence and transmission light channels.

Statistics

Data were visualized and analyzed using OriginLab software. Data groups were considered significantly different as indicated in the text, but considered not significantly different when P values were greater than 0.05 (ANOVA). Boxplots additionally show raw data points. Boxes indicate median, 25% quartile, and 75% quartile. Whiskers show minimal and maximal values. To plot the mean kinetics of reporter cleavage of single cell data, we set the time point that directly preceded cell death to time = 0 and calculated the mean and SD of single cell responses. The time values can, therefore, have a negative sign on the plots.

AUTHOR CONTRIBUTIONS

CL designed the study, performed experiments, analyzed the data, and wrote the paper. PS, IP, and DU performed experiments and analyzed data. MC isolated and purified primary human NK cells. RE, JB, and CW supervised the work and helped writing the paper.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at https://www.frontiersin.org/articles/10.3389/fimmu.2018.01840/full#supplementary-material.

FIGURE S1 | Transient knockdown of CD95 in HeLa cells. HeLa cells were non-transfected (black line) or transfected with a scrambled shRNA against CD95 (red line). Cells were stained 3 days after transfection with or without the anti-CD95 antibody DX2.

FIGURE S2 | Granzyme B and caspase-8 activity in HeLa and MDA-MB-468 cells upon killing by primary natural killer (NK) cells. (A) HeLa-CD48 and (B) MDA-MB-468 cells were transfected with NES-ELQTD-mGFP (for caspase-8) and NES-VGPD-mCherry for granzyme B and incubated with activated primary human NK cells. Reporter cleavage was analyzed as described in Figure 4.

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