Yersinia enterocolitica Targets Cells of the Innate and Adaptive Immune System by Injection of Yops in a Mouse Infection Model

Martin Köberle¹, Annegret Klein-Günther¹, Monika Schütz¹, Michaela Fritz¹, Susanne Berchtold¹, Eva Tolosa²,³, Ingo B. Autenrieth¹, Erwin Bohn¹*

¹Institut für Medizinische Mikrobiologie und Hygiene, Universitätsklinikum Tübingen, Tübingen, Germany, ²Hertie-Institut für klinische Hirnforschung, Universitätsklinikum Tübingen, Tübingen, Germany, ³Zentrum für Molekulare Neurobiologie, Universität Hamburg, Hamburg, Germany

Abstract

Yersinia enterocolitica (Ye) evades the immune system of the host by injection of Yersinia outer proteins (Yops) via a type three secretion system into host cells. In this study, a reporter system comprising a YopE-β-lactamase hybrid protein and a fluorescent staining sensitive to β-lactamase cleavage was used to track Yop injection in cell culture and in an experimental Ye mouse infection model. Experiments with GD25, GD25-β1A, and HeLa cells demonstrated that β1-integrins and RhoGTPases play a role for Yop injection. As demonstrated by injection of splenocyte suspensions in vitro, injection of Yops appears to occur randomly into all types of leukocytes. In contrast, upon infection of mice, Yop injection was detected in 13% of F4/80⁺, 11% of CD11c⁺, 7% of CD49b⁺, 5% of Gr1⁺, 2.3% of CD19⁺, and 2.6% of CD3⁺ cells. Taking the different abundance of these cell types in the spleen into account, the highest total number of Yop-injected cells represents B cells, particularly CD19⁺CD21⁺CD23⁺ follicular B cells, followed by neutrophils, dendritic cells, and macrophages, suggesting a distinct cellular tropism of Ye. Yop-injected B cells displayed a significantly increased expression of CD69 compared to non-Yop-injected B cells, indicating activation of these cells by Ye. Injection of IFN-γR (receptor-) and TNFRp55-deficient mice resulted in increased numbers of Yop-injected spleen cells for yet unknown reasons. The YopE-β-lactamase hybrid protein reporter system provides new insights into the modulation of host cell and immune responses by Ye Yops.

Introduction

Yersinia enterocolitica (Ye) is an enteropathogenic bacterium that causes gastrointestinal disorders such as enteritis and enterocolitis as well as extraintestinal manifestations such as lymphadenitis, reactive arthritis, and septicemia [1,2]. Ye has been demonstrated to multiply extracellularly in host tissue. To accomplish this, Ye needs to evade the host’s immune defense. Beside other virulence factors, Ye evolved a type III secretion system (TTSS) consisting of an injectisome and effector proteins the latter of which are injected into host cells [3]. The injection of effectors into host cells via a TTSS injectisome is a common strategy of pathogenic bacteria to counteract the host’s immune response [4]. The TTSS injectisome is complex ATP-driven protein-export machinery. Built of ring shaped proteins, the basal body is providing a channel through the bacterial membranes and the periplasm or the peptidoglycan wall, respectively. The injectisome is terminating in a needle-like structure that is protruding from the bacterial surface [5,6]. Thus, pore-forming proteins enable the injection of the effector proteins through the membrane of host target cells [7,8].

The TTSS is crucial for Yersinia virulence [9]. Ye injects at least six effector Yops into host cells. YopP/J is a potent inhibitor of the NF-κB and the MAPK signaling pathways and thus inhibits downstream effects of these pathways such as proinflammatory responses or antigen uptake [10–16]. In addition YopP induces apoptosis in macrophages and dendritic cells [17–20]. YopE, YopT and YopO affect RhoGTPase functions which leads to actin cytoskeleton disruption and together with YopH, a tyrosine phosphatase which targets different eukaryotic kinases, promote inhibition of phagocytosis [21]. In addition, phosphatase activity of YopH counteracts T cell activation [22,23]. YopM is known to interact directly with protein kinase C-like 2 (PRK2) and ribosomal S6 protein kinase 1 (RSK1), the function of YopM is so far elusive [24].

Despite well defined and profound in vitro effects of Yops, the contribution of some Yops to establish successful infection in a mouse infection model seems to be rather insignificant (YopP/O/T), while others (YopE/H/M) are of great importance for virulence of Ye [9]. This highlights the necessity to understand Yersinia pathogenicity in the context of whole organism infection. It also creates the need for a tool to display the sites of interaction between host organism and pathogen.

Various approaches have been taken to enable monitoring of type three secretion into eukaryotic cells by Yersinia and other pathogens during infection in cell culture. TTSS effectors have been detected in host cells by immunocytochemistry and western

Citation: Köberle M, Klein-Günther A, Schütz M, Fritz M, Berchtold S, et al. (2009) Yersinia enterocolitica Targets Cells of the Innate and Adaptive Immune System by Injection of Yops in a Mouse Infection Model. PLoS Pathog 5(8): e1000551. doi:10.1371/journal.ppat.1000551

Editor: Ralph R. Isberg, Tufts University School of Medicine, United States of America

Received October 15, 2008; Accepted July 22, 2009; Published August 14, 2009

Copyright: © 2009 Köberle et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: The project was partially funded by the Deutsche Forschungsgemeinschaft, SFB766(Bakterielle Zellhülle), BMBF/ERA-Net (RNAi-Net). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: erwin.bohn@med.uni-tuebingen.de
Author Summary

An important strategy of *Yersinia enterocolitica* (Ye) to suppress the immune defense is to inject bacterial proteins (*Yersinia* outer proteins, Yops) after cell contact directly into host cells, which affects their functions. However, tracking of cells in which Yop injection occurred has only been described for *Yersinia pestis* thus far. We adapted the described reporter system specifically for the use of infections with Ye and report the usefulness and limitations of this system. Using cell culture experiments, we demonstrated that β1-integrins and the RhoGTPases RhoA and Rac1 are involved in Yop injection. Since cell culture experiments also revealed that Yop injection is detectable in a similar manner into all subpopulations of the spleen, the system can be used to detect interaction of bacteria with host cells in vivo. In a mouse infection model we found that follicular B cells, granulocytes, macrophages, and dendritic cells are the main targets of Yop injection. Interestingly, Yop-injected B cells displayed an increased activation as indicated by increased CD69 expression. In contrast, interaction of bacteria with T cells seems to be rather a rare event. In immunocompromised gene-targeted mice we found increased frequencies of Yop-injected host cells for yet unknown reasons. Taken together, this novel reporter system represents a powerful tool to further study interaction of host cells with Ye.

cells and macrophages while injection into T and B cells was found to be a rare event. Geddes et al. [32] used translational fusions of bla and effectors of the SPI-1/2 pathogenicity islands as reporters and demonstrated that *Salmonella* targets preferentially granulocytes in the spleen after mouse infection.

To investigate into which immune cells Yops are injected during *Y. enterocolitica* infection in vivo, a Bla reporter system was established to track Yop injection in infection in cell culture and in an experimental mouse infection model. The established system was validated as a powerful tool to investigate Yop injection in cell culture as well as in a mouse infection model.

Results

Construction of a reporter system for tracking *Y. enterocolitica* Yop injection into host cells

To establish a reporter system to detect Yop injection by Ye, the expression vector pMK-Bla was transformed into E40 *Aad*, resulting in the strain E40-pBla, secreting a YopE53-β-lactamase fusion protein (see Materials and Methods as well as Table 1). As a control, a strain also harboring pMK-Bla but deficient in Yop secretion (ΔYscN-pBla) was employed. As a further control the plasmid pMK-Ova, which encodes for a translational fusion of YopE53 with the ovalbumin aa 247–355 fragment, was transformed into E40 *Aad* resulting in E40-pOva. Expression and secretion of YopE and the fusion proteins YopE53-Bla and YopE53-Ova by the strains E40-pBla, E40-pOva and ΔYscN-pBla was analyzed by immunoblotting. The data depicted in Figure 1A indicate that YopE wild type and hybrid proteins were expressed and secreted into the supernatant except by the ΔYscN-pBla mutant which is deficient for Yop secretion. To confirm injection of YopE and YopE53-Bla, HeLa cells were mock-infected or infected with E40-pBla or ΔYscN-pBla and injection assays were performed to detect YopE and YopE53-Bla by immunoblot using anti-YopE antibodies. YopE injection was detectable after infection with E40-pBla but not after infection with ΔYscN-pBla (Figure 1B left panel). In contrast, a YopE-Bla band was hardly visible. Only long overexposure led to a clearly visible YopE53-Bla band in E40-pBla, but not ΔYscN-pBla infected cells (Figure 1B right panel). This shows that YopE53-Bla is translocated into cells but with a much lower efficacy than the wildtype YopE.

Detection of Yop injection into epithelial cells in vitro

To investigate the usability of the system to assay Yop injection into host cells, HeLa cells were infected with the E40-pBla, E40-pOva or ΔYscN-pBla mutant strains and subsequently stained with CCF4-AM. In the presented study CCF4-AM was used instead of CCF2-AM that was utilized in former studies, because according to the manufacturer the FRET is slightly stronger with this dye, resulting in less background staining. Microscopic analysis of the uninfected cells as well as the controls including infection with E40-pOva or ΔYscN-pBla showed only green net fluorescence resulting from intact FRET within CCF4 (Figure 1C). In contrast, after E40-pBla infection, approximately half of the cells showed a pronounced blue net fluorescence resulting from FRET disruption by β-lactamase, suggesting that injection of the YopE53-Bla hybrid protein occurred (Figure 1C).

To quantify cell numbers and fluorescence signal intensity, flow cytometry was used. Green fluorescence resulting from CCF4-AM dye substrate uptake, de-esterification and retention in infected HeLa cells was equal to that observed in uninfected cells (Figure 1D). It also reflects the viability of cells because dead cells cannot retain the CCF4-dye (data not shown). After infection of HeLa cells for 1 hour with E40-pBla (MOI 50), blue fluorescence
To address whether reduced Yop injection into GD25 cells compared to GD25- cells is essential to detect Yop injection into host cells. Role of β1-integrins for Yop injection

Adhesion of Ye to host cells is mediated by interaction of host 1-integrins and the cell line GD25- 1-integrins are essential for Yop injection, infection of GD25 cells yielded 18.2% or 0.22% blue cells, respectively (Figure 2B). Infection of GD25 cells with E40-pBla or E40-pOva (MOI 50) for one hour resulted in 0.9% ± 0.2% or 1.2% ± 0.06% blue cells while infection of GD25-β1A cells yielded 18.2% ± 0.6% or 0.22% ± 0.12% blue cells, respectively (Figure 2B). The strong reduction (95%) of Yop injection into GD25 cells compared to GD25-β1A cells was also confirmed by detection of YopE by immunobLOTS after infection with E40-pBla (Figure 2C). To address whether reduced Yop injection into GD25 cells was due to reduced adhesion of Ye to GD25 compared to GD25-β1A cells, cell adhesion assays were performed. GD25 showed a modest (40%) but significant decrease of the number of adhering yersiniae compared to GD25-β1A cells (Figure 2D).

C. difficile toxin B (TcdB), an inhibitor of RhoA, was shown to inhibit Yop injection while the Rac inhibitor NSC23766 had no impact on Yop injection [35]. According to these data we hypothesized that RhoA might be crucial for Yop injection. To confirm the previous data, HeLa cells were pretreated with TcdB or NSC23766 and then infected with E40-pBla and stained with CCF4-AM (Figure 2E). Flow cytometry analysis revealed that the number of blue cells was significantly decreased by TcdB, but not by the Rac1 inhibitor NSC23766. Additionally, HeLa cells were transfected with control siRNA (siGo) or siRNAs specific for the RacGTPases Rac1, Cdc42 or Rac1. The cells were subsequently infected with E40-pBla, stained with CCF4-AM and analyzed by flow cytometry (Figure 2E). Inhibition of RhoA and Rac1, but not Cdc42, significantly reduced the number of blue cells by 25–30% indicating that both Rac1 and RhoA play a role in Yop injection.

Detection of Yop injection into leukocytes in vitro

To investigate whether Yops are also injected into primary cells, single cell suspensions were prepared from the spleen of C57BL/6

resulting from the cleavage of CCF4 was detectable in 58.9±11% of the viable cells. In contrast, infection of HeLa cells with E40-pOva or ΔYscN-pBla resulted in 0.48±0.3% or 2.5±2.4% blue cells, indicating that spontaneous cleavage of the substrate CCF4 after infection does not significantly occur and that Yop secretion is essential to detect Yop injection into host cells.

Table 1. Plasmids and bacterial strains used in this study.

| Designation       | Genotype or Description                                      | Reference or Source |
|-------------------|-------------------------------------------------------------|---------------------|
| pBME53-YopT       | pACYC184 derivative containing HindIII-sycE-yopE53-BamHI-yopT-Sall, CmR | [56]               |
| pBME53-Cre        | Derivative of pBME53-YopT, BamHI-yopT – Sall fragment was replaced by BamHI-nls-cre-Sall resulting in HindIII-sycE-yopE53-BamHI-nls-cre-Sall, CmR | This study         |
| pYV               | Cloning vector based on a small kryptic plasmid from Y. enterocolitica biogroup 1A strain 29807 modified with a kanamycin resistance marker and a pBluescript MCS. KanR | [57]               |
| pYV2-Syc-E-YopE53-Cre | Derivative of pYV2 with insertion of HindIII sycE-yopE53-BamHI-nls-cre-Sall into MCS. KanR | This study         |
| pMK4              | Derivative of pYV2-Syc-E-YopE53-Cre with insertion of asd under control of its own promoter between the Apol and Xbol sites. KanR | This study         |
| pMK1              | Sall-asd 3’UTR-EcoRI-asd 5’UTR- Xbol in pBluescript II SK(+). AmpR | This study         |
| pMK2              | Sall-asd 3’UTR-EcoRI-asd 5’UTR-Xbol in pMR5101. CmR         | This study         |
| pMK3              | asd mutator plasmid, derived from pMK2 by NotI digestion and relegation. CmR | This study         |
| pMR5101           | Mutator plasmid. CmR                                       | This study         |
| pBME53-Bla        | Derivative of pBME53-YopT. BamHI-yopT – Sall fragment was replaced by BamHI-bla-Sall resulting in HindIII sycE-yopE53-BamHI-bla-Sall, CmR | This study         |
| pMK-BlA           | HindIII sycE-yopE53-BamHI-NLS-Cre-Sall fragment of pMK4 was replaced by a HindIII sycE-yopE53-BamHI-bla-Sall fragment of pBME53-BlA. KanR | This study         |
| pYopE138-Ova147-355 | pACYC184 derivative containing HindIII-sycE-yopE138-BamHI-oVA147-355-Sall | [58]               |
| pBME53-Ova        | Derivative of pBME53-YopT. BamHI-yopT – Sall fragment was replaced by BamHI-oVA147-355-Sall, CmR | This study         |
| pMK-Ova           | Derivative of pMK-BlA. BamHI-bla-Sall was replaced by BamHI-oVA147-355-Sall from pBME53-Ova. KanR | This study         |
| pMSL41            | pYV40 with deletion in the yscN gene (yscN6169-177). ArsR | [27, 59]           |

Y. enterocolitica E40 strains

| E40               | Serotype O:9 patient isolate. NaI, ArsR | [27]               |
| E40 Δasd          | E40 strain with asd gene knockout, deficient in L-aspartate-dehydrogenase expression. NaI, ArsR, DBPMX | This study         |
| E40-pBla          | E40 Δasd strain transformed by pMK-BlA. NaI, ArsR, KanR | This study         |
| pYV Δasd pBla     | E40 Δasd pBla strain without pYV40 virulence plasmid. NaI, ArsR, KanR | This study         |
| ΔYscN-pBla        | pYV Δasd pBla strain transformed by pMSL41. NaI, ArsR, KanR | This study         |
| E40-pOva          | E40 Δasd strain transformed by pMK-Ova. NaI, ArsR, KanR | This study         |
Figure 1. Establishment of a β-lactamase reporter system and detection of Yop injection in HeLa cells. (A) Secreted (left panel) and intracellular (right panel) YopE, YopE53-Bla and YopE53-Ova hybrid proteins derived from supernatants or pellets, respectively, of bacterial cultures were detected by immunoblot using anti-YopE antibodies. To determine Yop injection into HeLa cells, cells were infected for 1 h (MOI 100) and injection of YopE hybrid proteins was detected (B) by immunoblot using anti-YopE and anti-actin antibodies. To visualize YopE53-Bla exposure time was extended (right panel). (C) Yop injection was detected as blue fluorescence by staining of cells with green fluorescent TEM-1 β-lactamase sensitive FRET substrate CCF4-AM and subsequent visualization by immunofluorescence microscopy. Blue and green fluorescence recordings were overlaid to get composite pictures. Blue and green cells are indicated with white and grey arrowheads, respectively. Alternatively, fluorescence of cells was detected by (D) flow cytometry. After flow cytometry data were analyzed and depicted as histograms. The upper panel shows uptake of CCF4-AM (green) by cells. The horizontal bar indicates the gating used to determine the number of β-lactamase positive cells revealing Yop-injection (blue) in the lower panel. Representative data of one of four experiments are shown. For note: percentages of blue or green fluorescent cells found in the histograms as depicted in this or other figures represent the results of this single experiment and are therefore discrepant to the means summarizing always several experiments as described in the results section.

doi:10.1371/journal.ppat.1000551.g001
Figure 2. Role of β1-integrins for Yop injection. GD25 and GD25-β1A cells were infected with E40-pOva or E40-pBla for one hour and subsequently stained with CCF4-AM and analyzed by flow cytometry. Living cells with high green fluorescence of CCF substrate were gated and are depicted in (A) dot blots. Data show the expression of β1-integrins (CD29) and blue fluorescence intensity of the coumarin product of uninfected or infected GD25 (upper panel) and GD25-β1A cells (lower panel) of one experiment. (B) Data show percentage of blue cells as the mean±SEM summarizing four independent experiments. Asterisks indicate significant differences (One-way ANOVA with Bonferroni corrections, p<0.001). (C) Injection of YopE into cells was detected by immunoblot using anti-YopE and anti-actin antibodies. (D) Adhesion assays were performed after infection of indicated cells with E40-pBla or E40-pOva for one hour. Bacteria and cells were stained with fuchsine and adherent bacteria per cell were counted. Data depict the mean and SD of two independent experiments. Hela cells were either (E) pretreated with TcdB (200 ng/ml) for 2 hours or pretreated with NSC23766 (100 μM) for three hours or (F) transfected with indicated siRNAs for 48 hours. Cells were subsequently infected with E40-pBla (MOI 50) for one hour and stained with CCF-AM and analyzed by flow cytometry. Data show percentage of blue cells as the mean±SEM of two (E) or three (F) independent experiments. Asterisks indicate significant differences compared to control (Paired t-test with control, p<0.05). doi:10.1371/journal.ppat.1000551.g002
mice and exposed to E40-pBla, E40-Ova or ΔYscN-pBla mutant strains at a MOI of 30 for one hour in vitro. Similar to infection of HeLa cells, flow cytometry analysis showed that CCF4 dye retention (green fluorescence) after infection of splenocytes was not significantly affected by infection with the strains used (data not shown). After infection of spleen cells for one hour with E40-pBla (MOI 50) in 61.4±11.4% of cells a blue fluorescence signal was detectable indicating that 61% of splenocytes had been injected with Yops (Figure 3A). In contrast, blue fluorescence was observed in less than 0.3% of splenocytes after infection with E40-pOva or ΔYscN-pBla. The low background of blue cells after infection with E40-pOva indicates that spontaneous cleavage of CCF4 detectable after infection with yersiniae is negligible. The low background staining after infection with ΔYscN-pBla indicates that secretion of YopE-Bla is a prerequisite for the detection of blue cells (Figure 3A) and that expression of YopE-Bla inside the bacteria and potential internalization of bacteria do not lead to significant blue fluorescence.

To determine the relationship between the number of bacteria per targeted host cell and the occurrence of blue cells (FRET disruption), spleen cell suspensions were infected with E40-pBla and E40-pOva as a control at various MOI for one hour and then subjected to flow cytometry analysis (Figure 3B and C). The data show that the percentage of blue cells correlates with the infection dose (MOI) in a fashion that a strong increase of the number of blue cells is followed by a saturation state. A hyperbolic regression curve can be calculated (goodness of fit of r² = 0.9977) with the formula: % blue cells = 84% MOI/(16.9 + MOI). Using this regression curve it can be calculated that an MOI of 16.9 would result in 50% blue cells. This regression curve also predicts that in cell culture even at high MOI Yop injection appears to be limited for yet unknown reasons.

We also investigated whether there might be preferential Yop injection into certain cell types. To define the populations of the spleen injected with YopE53-Bla, splenocytes were infected with E40-pBla and then flow cytometry analysis was used to detect distinct cell surface markers on and β-lactamase activity in the cells. The data depicted in Figure 4A shows the composition of spleen cells and confirmed that B cells (57±2% CD19⁺) and T cells (30±7% CD3⁺) are the most abundant spleen cell subpopulations. As shown in Figure 4B, the percentage of blue cells in each analyzed spleen cell population ranged between 58±6% to 82±10% but revealed no significant differences (p>0.05) indicating that Yop injection occurs into all cells types to a similar degree. Consistently, the composition of blue spleen cells (Figure 4C) is closely related to the frequency of the subpopulations.

Detection of Yop injection into spleen cells of Ye infected mice in vivo

To investigate whether Yop injection can be detected in vivo, desferrioxamine-conditioned C57BL/6 mice were intravenously infected with E40-pBla or E40-pOva; one to three days later splenocyte suspensions were prepared, stained with CCF4-AM and subjected to flow cytometry analysis (Figure 5A and Figure S1). As indicated in Figure 5A and Figure S1, after infection with E40-pBla a distinct population of blue cells (3.1±1.1% of total spleen cells) could be identified which was not present in the spleen of uninfected or E40-pOva (0.2±0.2%) infected mice. To confirm data obtained by flow cytometry, cell suspensions were subjected to flow cytometry cell sorting and subsequently analyzed by fluorescence microscopy (data not shown). Analysis of β-lactamase activity after infection for different time periods revealed a time-dependent increase of the percentage of blue cells with a maximum between two and three days after infection. The highest bacterial burden in the spleen was found between 24 and 48 hours after infection (Figure 5B). Increased infection dose led to a modest increase of the bacterial load in the spleen two days after infection (Figure 5C). However, an infection with more than 5×10⁸ E40-pBla did not further increase the bacterial burden in the spleen. The number of blue cells increased with the infection dose. In Figure 5D the bacterial load in the spleen and the percentage of blue cells from various experiments is given. The best fit to approximate the correlation of bacterial load and percentage of blue cells was obtained using a sigmoid regression curve (Figure 5D) (goodness of fit: r² = 0.64, degrees of freedom = 79, and Sy.x = 0.50). Plotting the percentage of blue cells versus the bacterial load in the spleen also revealed that for the detection of blue cells over background (defined as % blue cells±2-fold standard deviation = 0.6% blue cells) the bacterial load has to be more than log_{10} CFU 5.6 per spleen.

Increased activation of Omp-injected B cells

Splenocyte B cell populations comprise CD19⁺CD21⁺CD23⁻ (marginal zone B cells, MZ), CD19⁺CD21⁻CD23⁺ (follicular B cells, FO), and CD19⁺CD21⁺ (newly formed B cells, NFB) B cells. To address whether Yop injection occurs in all of these B cell subpopulations, cultured splenocytes were infected in vitro with E40-pBla (MOI 50) for 1 hour and Yop injection was determined after staining using antibodies for B cell markers and CCF4 (Figure S2). The data show ~84% of B cells were blue indicating injection of Yops; moreover, the different B cell subpopulations contributed to the total number of CD19⁺ blue cells according to their different frequencies of MZ (~10%), FO (~70%) and NFB (~20%), as injection of Yops occurred with similar efficacy in all B cell subtypes in vitro.

To analyze Yop injection into B cells in vivo, desferrioxamine-conditioned C57BL/6 mice were infected with 5×10⁸ E40-pBla and two days later B cells were analyzed (Figure 6A). After
Figure 3. Detection of Yop injection in cultured splenocytes. Splenocytes were infected with (A) different Yersinia strains (MOI 50) or (B) with different infection doses of E40-pBla for 1 h. Subsequently cells were stained with CCF4-AM (A–B) and flow cytometry analysis was performed. Data in A, depicted as histograms, show the green fluorescence of the substrate CCF4 and the percentage of green cells (upper panel) is marked. Only these cells were gated to determine the number of blue fluorescent cells indicating injection of effector Yops and the β-lactamase reporter (lower panel). Data in B depict the percentage of blue fluorescent cells after infection with E40-pBla or E40-pOva using different MOI and these data are summarized in (C) for infection with E40 as mean±SEM of four independent experiments. The dotted line represents a calculated hyperbolic regression curve (goodness of fit: r² = 0.977).

doi:10.1371/journal.ppat.1000551.g003
infection, the composition of splenic B cell subpopulations changed. Thus, the percentage of CD21\(^+\)CD23\(^-\) (FO) (43.8±4 in infected versus 66%±2 in uninfected mice) and the percentage of CD21\(^+\)CD23\(^-\) (MZ) (3.6±0.9 in infected versus 9.2±1.4 in uninfected mice) significantly decreased whereas the percentage of CD21\(^-\)NFB (47.2±3.8 in infected versus 22.2±0.9 in uninfected mice) significantly increased. Taking the changes in total cell numbers in the spleen into account, the total number of NFB significantly increased while the total number of FO and MZ remained largely unchanged (data not shown). Analysis of blue (Yop-injected) B cells (~2% of total B cells) revealed that 64.4%±1.7 of CD19\(^+\) blue B cells were CD21\(^+\)CD23\(^-\) (FO) B cells indicating that *Yersinia* targets predominantly follicular B cells (Figure 6C). In blue FO B cells, a distinct CD21\(^+\)CD23\(^-\) B cell subpopulation (Figure 6D) was detected suggesting that injection of Yops into FO is associated with an increased expression of CD21 and CD23.

To address whether injection of Yops may affect activation of B cells, expression of CD69, an early activation marker of B cells, was determined. In splenocytes infected in vitro with E40-pBla, increased CD69 expression was found in B cells (mean fluorescence intensity, MFI; 7.5 in uninfected versus 29.8 in infected cultures). In splenocytes infected in vitro with E40-pBla, increased CD69 expression was found in B cells (mean fluorescence intensity, MFI; 7.5 in uninfected versus 29.8 in infected cultures).

Upon infection of mice with E40-pBla, ~2% of B cells were infected with Yops (blue staining) (Figure 7B). Expression of CD69 was increased in splenic B cells compared with uninfected mice (MFI of 26.1 in infected versus 12.3 in uninfected mice). However, in Yop-injected B cells (green\(^+\) blue\(^-\)), expression of CD69 was higher compared with B cells not infected with Yops (green\(^+\) blue\(^-\)) (Figure 7A) suggesting that in vitro, injection of Yops into B cells is associated with rapid activation of these cells.

Infection of TNF-Rp55 and IFN-γR deficient mice

Previous studies showed that TNFRp55\(^-\)/ and IFN-γR\(^-\)/ mice are highly susceptible to infection with Ye [36–40] indicating that the pleiotropic effects of IFN-γ and TNF-α are crucial for the defense against *yersinia*. To investigate whether such gene defects may modulate Yop injection, desferrioxamine conditioned TNFRp55\(^-\)/, IFN-γR\(^-\)/, and wild type mice were infected with Ye and two days later, splenocytes suspensions were prepared, stained with antibodies and CCF4-AM, and subjected to flow cytometry analysis. Infection of TNFRp55\(^-\)/ and IFN-γR\(^-\)/ mice with E40-pOva as well as E40-pBla resulted in a bacterial load of log\(_{10}\)CFU 7.4±0.1 and 7.3±0.1, respectively and was comparable to the bacterial load of infected WT mice (log\(_{10}\)CFU 7.4±0.1) (Figure 8A).

After infection of IFN-γR\(^-\)/ mice no significant changes in the splenic cell subpopulations were observed (Figure 8C) compared to infection of WT mice. However, the percentage of all blue cells increased significantly (IFN-γR\(^-\)/ 5.9±0.9% versus WT 3.1±1.1%) (Figure 8B). Likewise, the percentage of blue Gr-1\(^+\), F4/80\(^+\), CD11c\(^+\) and CD49b\(^+\), but not CD3\(^+\) and CD19\(^+\), cells increased significantly (Figure 8D). Taking both, the total number of blue cells as well as changes in the cell composition into account, no significant alterations in the composition of blue cells was found (Figure 8E). These data indicate that IFN-γR deficiency leads to an increased number of cells, in which Yops are injected, but the composition of blue cells is comparable with that of infected C57BL/6 mice.

Infection of TNF-Rp55\(^-\)/ mice led to a significant increase in the percentage of Gr-1\(^+\), F4/80\(^+\) and CD11c\(^+\) cells, and a significant decrease of C19\(^+\) cells compared with WT mice (Figure 8C). The percentage of total blue cells was significantly higher after infection of TNF-Rp55\(^-\)/ (7.6±1.1%, p<0.001) compared with infection of WT mice (3.1±1.1%) (Figure 8B). In line with these findings, all subpopulations (F4/80, Gr-1, CD11c, CD49b, CD19 and CD3) showed a significantly increased percentage of blue cells in infected TNFRp55\(^-\)/ mice (Figure 8D). The composition of blue cells changed leading to a significantly increased number of blue Gr-1\(^+\) and F4/80\(^+\) cells compared to WT mice (Figure 8E). These data suggest that TNFRp55 deficiency leads to a more frequent injection of Yops into F4/80\(^+\) and Gr-1\(^+\) cells.

To investigate the localization of *yersinia* in the spleen after infection of mice, splenic sections were prepared and immunohistochemistry using anti-*Yersinia* Hsp60 antibodies was performed (Figure 8F). Histological and immunohistochemical studies revealed that Ye formed...
Figure 5. Detection of Yop injection in splenocytes derived from infected mice. Mice were infected with indicated Yersinia strains and isolated splenocytes were stained with CCF4-AM. In some experiments, cells were additionally stained with antibodies specific for cell surface markers (B–D). Subsequently cells were analyzed by flow cytometry. (A) Dot plots show cells gated on high levels of green fluorescence of the substrate CCF4 (viable cells) plotted versus the blue fluorescence revealing Yop-injection (infection dose $5 \times 10^5$ for 2 days). (B) Four mice (two independent experiments with each two animals) per group were infected with $5 \times 10^5$ E40-pBla for indicated time periods or with (C) indicated numbers of bacteria for two days and the log$_{10}$ CFU in the spleen (upper panel) and the percentage of blue cells (lower panel) was determined. Data represent the mean±SD. (D) summarizes the correlation between the log$_{10}$ CFU and the percentage of blue cells in the spleen of E40-pBla infected C57BL/6 mice by using data of experiments with different infection dose as well as different time ranges of infection (n = 84). A sigmoidal regression curve was calculated (goodness of fit $r^2 = 0.64$, sy.x = 0.5). (E) summarizes the percentage of splenocytes expressing the indicated surface markers prior to and after infection with indicated Yersinia strains (infection dose $5 \times 10^5$ for 2 days). Data represent the mean±SD of 5 uninfected, 5 E40-pOva or 22 E40-pBla infected mice of more than five independent experiments. (F) depicts the percentage of blue cells after infection with E40-pBla for each indicated subpopulation and (G) shows the mean±SD of the percentage of blue cells expressing one of the indicated surface markers summarizing 11 independent experiments with two mice (lowest panel), n = 22. Comparison of all groups with each other by one-way ANOVA analysis and Bonferroni correction revealed significant differences p<0.05 in: (B) for CFU and for % blue cells as indicated by asterisks, in (C) for CFU: $5 \times 10^1$ versus $5 \times 10^3, 5 \times 10^4$, or $5 \times 10^5$ and $5 \times 10^2$ versus $5 \times 10^3, 5 \times 10^4$, and $5 \times 10^5$, for % blue cells: $5 \times 10^1, 5 \times 10^2$ or $5 \times 10^3$ versus $5 \times 10^4, 5 \times 10^5$, or $5 \times 10^6$, $5 \times 10^4$ versus $5 \times 10^5$. Asterisks indicate differences in (E) if uninfected versus infected is compared (One-Way ANOVA with Dunnett corrections, p<0.05).

doi:10.1371/journal.ppat.1000551.g005
many microcolonies surrounding the lymphoid follicles. Gross differences in the splenic architecture or localization of yersiniae in the spleen of WT or knockout mice were not observed.

To further address why in TNFRp55^−/− and IFN-γR^−/− mice increased numbers of Yop injected spleen cells were found, splenocytes of C57BL/6, TNFRp55^−/− and IFN-γR^−/− mice were infected in vitro for one hour with different MOI of E40-pBla and the percentage of blue cells was determined. The frequency of blue cells for the different MOIs was not significantly different between C57BL/6 and TNFRp55^−/− or IFN-γR^−/− splenocytes (Figure S4) suggesting that splenocytes of wildtype, TNFRp55^−/− and IFN-γR^−/− mice upon infection with Ye display a comparable frequency of Yop injection.

**Discussion**

Injection of Yop effector molecules into host cells by the TTSS is an important strategy of Ye to suppress the host immune response. However, detection of Yop injection in animal infection models has been difficult so far. Several tools have been employed to detect injection of bacterial proteins into host cells, but their usefulness has been limited to certain in vitro models [8,25–28]. In similar, results published by Briones et al. and own attempts to create a Cre-recombinase based reporter system to monitor Yop injection into host cells produced promising results in cell culture experiments but failed to deliver a valuable tool in mouse infection models [29]. Recently however, Marketon et al. demonstrated that Yop-β-lactamase fusion proteins can be utilized to monitor Yop injection in vivo [30]. In this study we applied this tool to Ye infection and tested whether it might be useful to detect Yop injection in vitro and in vivo.

In line with the report by Marketon et al. [30], the use of control strains such as ΔYscN-pBla and E40-pOva revealed that detection of β-lactamase reporter activity depends on injection of the YopE53-Blal fusion protein. Therefore, internalization of bacteria does not account for reporter activity. Yop injection into host cells...
Figure 7. Changes of CD69 expression in splenocyte populations after *Yersinia* infection. (A) Cultured splenocytes were infected with E40-pBla (MOI 50) or mock-infected for 1 hour or (B) desferrioxamine conditioned C57BL/6 mice were left untreated or infected with $5 \times 10^5$ E40-pBla for two days and the splenocytes were then isolated. Splenocytes were stained with anti-CD19-APC, anti-CD69-PE-Cy7 and subsequently with CCF4 and analyzed by flow cytometry. Cells were gated (as shown in Figure S3) for high green fluorescence (viable cells) and B cells (CD19$^+$). Left panels show histograms for CD69 expression of non-infected cells gated for viable CD19$^+$ cells (black line) or of E40-pBla infected cells gated for all viable CD19$^+$ cells (total, red line), gated for green$^+$ blue$^+$ CD19$^+$ cells (blue line) or green$^+$ blue$^2$ CD19$^+$ cells (green line). Mean fluorescence intensities (MFI) of CD69 expression are depicted for this representative experiment. Right panels show the means and SEM of the x-fold increase of MFI of the indicated cells compared to uninfected of (A) three independent experiments or (B) two independent experiments with a total of five mice per group. Asterisks indicate significant differences, $p < 0.01$.

doi:10.1371/journal.ppat.1000551.g007
Figure 8. Impact of IFN-γR and TNF-R deficiency on Yop injection. IFN-γR<sup>−/−</sup>, TNFRp55<sup>−/−</sup> and control C57BL/6 mice were infected with E40-pBlu and two days later (A) bacterial load in the spleen and (B) percentage of blue cells was determined. Each dot depicts the results for one single mouse (in total 8–22 mice were investigated). Data in (C) show the mean and SD of the distribution of cell subpopulations expressing indicated markers, in (D) the mean and SD of percentage of blue cells of each indicated subpopulation and in (E) the mean and SD of the percentage of the indicated subpopulations of blue cells. Asterisks show significant differences between all groups compared (A, B, one-way ANOVA with Bonferroni corrections, p<0.05) and differences between control mice and knockout mice (C, D, E; one-way ANOVA, with Dunnett corrections, p<0.05). F. Immunohistology of infected spleen of indicated mice. Splenic sections were stained with anti-hsp60 and counterstained with Mayer's hemalaun. White arrowheads mark lymph follicle and black arrows colonies of yersiniae. All the experiments were repeated more than four times using in total 8–22 mice. Marked regions are shown at higher magnification in lower panels.
doi:10.1371/journal.ppat.1000551.g008
was detectable after infection of HeLa epithelial cells and primary C57BL/6 splenocytes in cell culture experiments. In vitro experiments with splenic subpopulations indicated that Yop injection occurs with similar efficacy into macrophages, dendritic cells, granulocytes, NK cells, T cells or B cells in cell culture. We conclude that the different immune cell subpopulations display the same prerequisites for Yop injection. From these data it is conceivable that Yop injection in vivo actually reflects the interaction frequency of bacteria with distinct splenic subpopulations.

The usefulness of this reporter system in cell culture experiments could be demonstrated by investigating the importance of β1-integrins for Yop injection. It is widely accepted that β1-integrins play an important role for adhesion to host cells and eventually for internalization of yersiniae via the direct or indirect interaction with invasin and YadA, respectively [34,41–43]. In line with this, the number of adhering Ye to fibroblasts which lack β1-integrins was reduced by 40% compared to cells expressing β1-integrins. In contrast, Yop injection (measured as percentage of blue fluorescent cells) into fibroblasts which lack β1-integrins was dramatically reduced by 95% compared to fibroblasts expressing β1-integrins. These data are in line with a recent report [35] which suggested a central role for β1-integrins for Yop injection. Nevertheless, detection of YopE translocation by immunoblot was still detectable in GD25 cells which lack β1-integrins, indicating that other surface molecules beside β1-integrins may also play a role to modulate Yop injection. At this point it is not clear which other surface molecules beside β1-integrins affect Yop injection and whether depending on the cell types in which Yops are injected different surface molecules modulate Yop injection in vitro and in vivo.

After *Y. pseudotuberculosis* infection β1-integrin mediated signal transduction leading to RhoGTPase activation seems to be the prerequisite for initiation of Yop effector injection [35]. By using the Rac1 inhibitor NSC23766 it was excluded that Rac1 plays a role for Yop injection and because both TcdB (inhibitor of RhoA, Rac1, Cdc42, RhG, Tec10) and C3 exotoxin (RhoA, RhoB, RhoC inhibitor) inhibited Yop injection, RhoA, RhoB and RhoC were identified as putative candidates for this effect [35]. In contrast, the siRNA experiments presented herein suggest that both Rac1 and RhoA play a role for Yop injection after Ye infection of HeLa cells. Interestingly, in the present study the Rac1 inhibitor NSC23766 also did not inhibit Yop injection. NSC23766 was demonstrated to bind to Rac1 and to prevent binding to and activation of Rac1 by the Rac1-specific guanine nucleotide exchange factors (GEF) Tiam1 and Trio. However, it was described that NSC23766 seems not to prevent binding to and activation of Rac1 by the more promiscuous GEF Vav [44,45]. As Vav is able to activate both RhoA and Rac1, but not Cdc42 [44,45], one may speculate that Vav or other GEFS, rather than Tiam1 and Trio, might be involved in Rac1 mediated Yop injection. One has to anticipate that NSC23766 inhibits *Yersinia* triggered Rac1 activation only partially and in a GEF specific manner which has to be proven in future studies. In addition it will be interesting to find out which specific GEFS are involved in facilitating Yop injection. Taken together this study clearly pinpoints that β1-integrins play a role in facilitating Yop injection.

The YopE-Bla reporter system turned out to work highly efficiently when expressed in E40 (O9) strain, but not in WA-314 (O8) strain; thus, 80% of in vitro infected splenocytes displayed blue fluorescence upon infection with E40-pBla strain, but only 26% upon infection with WA-pBla strain (data not shown). Moreover, in C57BL/6 mice infected with E40-pBla 3% of splenocytes displayed blue fluorescence, while in WA-pBla infected mice we could hardly detect any blue cells for yet unknown reasons. We therefore used the E40 strain for all experiments included in this study. The low virulence of serotype O9 strains such as E40 strain or O3 strains in mice is at least partially caused by their disability to synthesize the iron chelating siderophore yersiniabactin due to their lack of the HPI island [46,47]. This could be “complemented” by treatment of mice with desferrioxamine which allows to study Ye O9 and Ye O3 infection in mice [48–50]. The disadvantage of desferrioxamine, however, is that it might have immunosuppressive effects on phagocytes [51,52].

Infection of desferrioxamine-conditioned C57BL/6 mice revealed a correlation between the percentage of blue cells and the bacterial burden in the spleen which can be best described by a sigmoidal dose response curve. Taking the background staining of 0.2±0.2% into account, a detection limit of 0.6% blue cells was estimated using the sigmoidal regression curve. To reach 0.6% blue cells in the spleen the bacterial burden would have to be at least 4×10^7 bacteria representing a MOI of 0.01. The cell culture experiments, however, showed that much higher MOI are required to achieve half-maximal numbers of blue spleen cells. Therefore, higher efficacy of YopE-Bla detection might be due to abscess formation with Ye microcolonies which may allow more Ye bacteria to interact with single host cells at abscesses. Alternatively the TTSS machinery might be upregulated during infection in spleen compared to cell culture. Most experiments were carried out at conditions with a bacterial burden of 5×10^7 bacteria in the spleen resulting in 3.1±1.1% of blue, Yop-injected cells.

Determination of the percentage of blue cells in each subpopulation revealed that Yop injection occurred into Gr1+ (predominantly granulocytes), CD11c+ (predominantly dendritic cells), F4/80+ (predominantly macrophages) and CD49b+ cells (predominantly NK cells). In contrast, only in a small percentage of the CD3+ (T cells) and CD19+ (B cells) cells, Yop injection was detectable. However, due to the high total number of B cells (>50% of spleen cells) their contribution to the total number of blue cells is about 40%. Further characterization of B cells revealed that the composition of B cell subpopulations changed after infection leading to an increased number of CD19+CD21hi (NFB) while the number of CD19+CD23+CD21lo cells (FO) did not change and the number of CD19+CD21hi CD23- (MZ) slightly decreased. Thus, after infection NFB appear to either proliferate in or immigrate into the spleen.

Yop injection was predominantly detectable in follicular B cells and to a lesser extent in NFB. Naïve follicular B cells reside in the “follicular niche” and may present T-dependent antigens to activated T cells. The follicular niche therefore represents the major site at which recirculating B cells mediate T-dependent immune responses to protein antigens [53]. Interestingly, in contrast to the Ye infection model presented here, injection of the Bla-reporter into B cells was not described in *F. peitidis* infection [30]. It remains unclear whether these discrepancies are due to different *Yersinia* species, different mouse strains or differences in the reporter systems. Nevertheless, it is striking that after *F. peitidis* infection the analysis gate which was used to define the viable cells was rather small indicating that most spleen cells were necrotic or apoptotic under the chosen experimental conditions. It might well be that Yop-injected B cells might have been missed by the analysis due to high cell death rates in *F. peitidis* infection.

In Ye infection, however, interaction of B cells with Ye is prominent. This finding is in line with the analysis of histological sections of the spleen. Thus, E40-pBla microcolonies were observed adjacent to lymphoid follicles (B cell zones). Balada-
Llasat and Mecsas [54] reported that Y. pseudotuberculosis may show a tropism to B and T cell zones in lymph nodes. In our studies we found yersiniae always adjacent to lymph follicles in the spleen and the most frequent interaction of yersiniae with host cells was that with B cells, specifically with CD19+/CD21+/CD23+ B cells which are defined as follicular B cells (FO). FO B cells organize into the primary follicles of B cell zones located around follicular dendritic cells in the white pulp of the spleen. Thus in our system Ye appear to preferentially colonize in the follicular niche and interact frequently with follicular B cells. Whether Ye are actively migrating to this region or may be trapped by e.g. dendritic cells needs to be further investigated. Infection with Ye was associated with increased CD21 and CD23 expression levels of FO B cells which suggests that B cells may be activated after interaction with yersiniae. Specific activation of Yop injected B cells was also supported by the observation that the early B cell activation marker CD69 was upregulated in blue B cells. From this data we can conclude that Yop-injected B cells were specifically activated by Ye despite injection of Yops. Whether Yop-injected B cells may undergo subsequent cell death or are affected in their ability to produce IgM or to undergo differentiation needs to be investigated in future study.

Previous studies revealed that TNF-α and IFN-γ are important for defense against Ye infection [36–38,40]. Therefore, we were interested whether these cytokines may affect Yop injection. The experimental conditions were chosen in a way that the bacterial burden in the spleen of the various mouse strains was comparable. Infection of IFN-γR- and TNFRIp55-deficient mice revealed a higher number of blue cells in the spleen compared to infected WT mice indicating that either more host cells interacted with yersiniae or the threshold which allows detection of Yop infection was lower in knock-out mice. Histological analysis of the Ye microcolonies in WT and knockout mice revealed comparable tissue distribution of Ye and comparable tissue alterations by the infection. In vitro infection of splenocytes from IFN-γR- and TNFRIp55 deficient and wildtype mice did not reveal significant differences in terms of Yop injection. Therefore it remains widely unclear why more blue cells are found in IFN-γR- or TNFRIp55−/− mice compared to wild type mice. However, we cannot exclude that minor changes in spleen cell subpopulations may partially cause the different numbers in knock-out and wild type mice.

In further experiments we tried to detect Yop injection into cells of the PP after orogastric infection. However, we were not able to detect blue PP cells under these conditions (data not shown). One explanation might be that the number of bacteria that establish infection in the PP is lower than that in spleen [55]. Thus, one may speculate that the number of bacteria attached to a single host cell in PP might be lower than that in spleen which might result in Yop injection below the detection limit of this reporter system. Alternatively or in addition, the number of cells in which Yops were translocated might be too low.

The Bla reporter system described herein is so far the only system that allows tracking of Yop injection in Ye infection in vivo on a single cell level in a mouse infection model. Nevertheless, the system needs to be improved in order to accomplish a more sensitive reporter system. The disadvantages of CCF-4 as a substrate for β-lactamase is that the amount of Yops translocated cannot be quantified and that CCF-4 displays limited sensitivity. Moreover, CCF-4 does not allow studying Yop injection in histological sections in situ. Current experiments in our laboratory aim at overcoming these limitations with new compounds.

Taken together, the Ye Bla reporter system can be successfully used in cell culture and in mouse infection models for detection of Yop injection on a single cell level, and thus could be a valuable tool for quantitatively screening for genetic factors involved in Yop secretion and injection at the bacterial as well as at the host cell side, or to identify drugs which target Yop secretion or injection, respectively.

Materials and Methods

Bacterial strains and growth conditions

Y. enterocolitica E40 strains used in this study are listed in table 1. All bacterial strains were grown overnignt in Luria-Bertani (LB) broth at 27°C supplemented with 50 μg/ml meso-diaminopimelic acid, 50 μg/ml nalidixic acid, 50 μg/ml kanamycin, and 400 μM Na-arsenate (all from Sigma Chemical, St. Louis, MO) in combinations according to the indicated resistances and supplementation needs (table 1). A 1:25 dilution of the bacterial culture was incubated for additional 3 h at 37°C. The bacteria were washed once with phosphate-buffered saline (PBS; Invitrogen, Karlsruhe, Germany) and the optical density at 600 nm was determined.

Plasmid constructions

All plasmid constructs and bacteria used are listed in table 1. To generate pMK-Bla, as a starting vector pBME53-yopT [56], a pACYC184 derive containing a HindIII-sycE-yopE/sycE promoter-yopE gene fragment encoding the first 56 N-terminal amino acids (aa) - BamHI-YopT - SalI construct was used. YopT was replaced by a translational fusion of the coding sequences (CDS) of the SV40 nuclear localization signal (nl) and the CDS of Cre recombinase resulting in pBME53-Cre. This vector was digested with HindIII and SalI and the resulting sycE-yopE53-cw fragment was inserted into pIV2 [57]. pIV2 was derived from a small cryptic plasmid of an apathogenic Y. enterocolitica strain (kindly provided by EckhardStrauch, Federal Institute for Risk Assessment, Berlin) resulting in pIV2-SycE-YopE53-Cre. To create a balanced system that compels stable retention of the reporter vector in Y. enterocolitica during in vivo infections, the Y. enterocolitica asd gene coding for the L-aspartate-dehydrogenase, an enzyme that is required for the synthesis of the cell wall component L-lysine via the meso-2,6-diaminopimelic acid (DAP) pathway and therefore essential for bacterial growth and replication was inserted into the pIV2-SycE-YopE53-Cre plasmid as follows: The asd gene and its promoter region were amplified from Y. enterocolitica E40 chromosome by using the oligonucleotides 5′-AGC TTT AGG GCC CAA AAA CAG CAA CAC CGT TGC C-3′ and 5′-AAC TCG ATC AAC TCG AGT TAC AGA AAA TTC GCA GC-3′. The PCR product was then digested with Apal and Xhol and inserted into those sites of pIV2-SycE-YopE53-Cre, thus yielding pMK4. Together with the deletion of the asd gene from its chromosomal locus (see below) in the utilized bacteria, this grants stable retention of the reporter plasmid, by complementing the otherwise lethal DAP auxotrophy of the Y. enterocolitica E40 Aasd strain.

The β-lactamase (bla) gene was amplified from pCR2.1 [Invitrogen, Karlsruhe, Germany] using the primers 5′-GGA TAC ATG AGT ATT CAA CAT TTC GC-3′ and 5′-GTC GAC AAC TTG GTC GTA CAG TTA CC-3′. The PCR product was subcloned into pCR-Blunt II and subsequently donated as a BamHI / SalI fragment to pBME53-YopT, thus replacing the sequence coding for YopT and fusing the translation product to the YopE53 domain, yielding the plasmid pBME53-Blal. pBME53-Bla was digested with HindIII-SalI and the resulting HindIII-sycE-yopE53-bla-SalI fragment replaced the HindIII-sycE-yopE53-cw-SalI fragment of pMK4 yielding pMK-Bla.

To generate a control vector, the sequence coding for β-lactamase was excised by BamHI / SalI digestion from pBME53-bla and
The plasmid pMSL41 has been yielded by deletion of the serotype O:9 [27]. To generate Y. enterocolitica isolate that belongs to the serotype O:9 [27,61] into pYV2 previously [62]. Overnight cultures grown at 27°C were diluted 1:20 into brain heart infusion broth (Difco, Heidelberg, Germany). 3 h after induction secreted proteins were examined by the preparation of released proteins as described above. Protein concentration was determined by Bradford assay (BioRad, Hercules, USA) and the 32P incorporation was extended to 30 min for increased sensitivity. One hour after infection (MOI 50), cells were washed with PBS, fixed with 4% PFA and stained with fuchsine. All samples were prepared in duplicates, of which cells and bacteria were counted in 6 representative details.

Analyses of Yop secretion

Secretion of the YopE53β-lactamase fusion protein was examined by the preparation of released proteins as described previously [62]. Overnight cultures grown at 27°C were diluted 1:20 into brain heart infusion broth (Difco, Heidelberg, Germany). After 2 h of incubation at 37°C, Yop secretion was induced by the addition of 5 mM EGTA for Ca2+ sequestration, 15 mM MgCl2 and 0.2% glucose. 3 h after induction secreted proteins were precipitated from the culture supernatant with trichloroacetic acid. Protein concentration was determined by Bradford assay (BioRad, Hercules, USA) and 15 μg of protein was loaded on a 12% SDS-PAGE. Separated proteins were transferred electrophoretically to an Immobilon-P PVDF membrane (Millipore, Bedford, USA). For immunostaining, polyclonal rabbit anti-YopE was used. Immunoreactive bands were visualized by incubation with peroxidase-conjugated swine anti-rabbit IgG antibody (1:1000) (DAKO, Glostrup, Denmark) using enhanced chemiluminescence reagents (ECL, Amersham Biosciences, Freiburg, Germany) and CL-XPosure Film (Thermo, Rockford, IL).

Cell culture and in vitro infections

The fibroblast-like GD25 and GD25-B1A cell lines were a kind gift from R. Fassler (Max-Planck-Institute for Biochemistry, Martinsried, Germany). The murine GD25 cells were derived from the embryonic stem cell clone G201 which is deficient in the β1-integrin subunit [63]. The stably transformed cell line GD25-B1A was obtained by electroporation of wild-type integrin β1A cDNA into GD25 cells [64]. GD25 cells not expressing β1-integrins were cultivated in DMEM (Gibco) supplemented with 10% FCS, the GD25-B1A line expressing β1-integrins was maintained in DMEM supplemented with 10% FCS and 10 μg/ml puromycin.

HeLa cervical epithelial cells (ATCC CCL-2.1) were grown in RPMI 1640 (Biochrom KG, Berlin, Germany) supplemented with 10% fetal bovine serum (Sigma Chemical), 2 mM L-glutamine (Biochrom KG), penicillin (100 U/ml) and streptomycin (100 μg/ml) (Biochrom KG) in a humidified 5% CO2 atmosphere at 37°C.

To enable the generation of reporter strains with pYV40 virulence plasmids that had been subjected to mutagenesis, the pMK3 asd mutator plasmid was constructed by allelic exchange. The pMK3 asd mutator plasmid was constructed by electroporation of pMSL41 [27,61] into pYV2 previously [62]. Overnight cultures grown at 27°C were diluted 1:20 into brain heart infusion broth (Difco, Heidelberg, Germany). 3 h after induction secreted proteins were examined by the preparation of released proteins as described above. Protein concentration was determined by Bradford assay (BioRad, Hercules, USA) and the 32P incorporation was extended to 30 min for increased sensitivity. One hour after infection (MOI 50), cells were washed with PBS, fixed with 4% PFA and stained with fuchsine. All samples were prepared in duplicates, of which cells and bacteria were counted in 6 representative details.

Analyses of Yop secretion

Secretion of the YopE53β-lactamase fusion protein was examined by the preparation of released proteins as described previously [62]. Overnight cultures grown at 27°C were diluted 1:20 into brain heart infusion broth (Difco, Heidelberg, Germany). After 2 h of incubation at 37°C, Yop secretion was induced by the addition of 5 mM EGTA for Ca2+ sequestration, 15 mM MgCl2 and 0.2% glucose. 3 h after induction secreted proteins were precipitated from the culture supernatant with trichloroacetic acid. Protein concentration was determined by Bradford assay (BioRad, Hercules, USA) and 15 μg of protein was loaded on a 12% SDS-PAGE. Separated proteins were transferred electrophoretically to an Immobilon-P PVDF membrane (Millipore, Bedford, USA). For immunostaining, polyclonal rabbit anti-YopE was used. Immunoreactive bands were visualized by incubation with peroxidase-conjugated swine anti-rabbit IgG antibody (1:1000) (DAKO, Glostrup, Denmark) using enhanced chemiluminescence reagents (ECL, Amersham Biosciences, Freiburg, Germany) and CL-XPosure Film (Thermo, Rockford, IL).

Cell culture and in vitro infections

The fibroblast-like GD25 and GD25-B1A cell lines were a kind gift from R. Fassler (Max-Planck-Institute for Biochemistry, Martinsried, Germany). The murine GD25 cells were derived from the embryonic stem cell clone G201 which is deficient in the β1-integrin subunit [63]. The stably transformed cell line GD25-B1A was obtained by electroporation of wild-type integrin β1A cDNA into GD25 cells [64]. GD25 cells not expressing β1-integrins were cultivated in DMEM (Gibco) supplemented with 10% FCS, the GD25-B1A line expressing β1-integrins was maintained in DMEM supplemented with 10% FCS and 10 μg/ml puromycin.

HeLa cervical epithelial cells (ATCC CCL-2.1) were grown in RPMI 1640 (Biochrom KG, Berlin, Germany) supplemented with 10% fetal bovine serum (Sigma Chemical), 2 mM L-glutamine (Biochrom KG), penicillin (100 U/ml) and streptomycin (100 μg/ml) (Biochrom KG) in a humidified 5% CO2 atmosphere at 37°C. Suspensions of single spleen cells were prepared from naïve C57BL/6 mice as described below. For infection with a MOI of 100 for microscopic observation and Western Blot analysis or with a MOI of 50 or as indicated for FACS analysis bacteria were spun onto the cells (5 minutes, 400 g). After 1 h incubation at 37°C, gentamicin (100 μg/ml) was added to terminate infection. To assess the impact of inhibitors of RhoGTPases on Yop injection, cells were treated with 200 ng/ml medium Clostridium difficile toxin B 10463 (TcdB; kind gift I. Just, Institute for Toxicology, Hannover Medical School, Hannover, Germany) 2 h prior to infection or 100 μM Rac1 inhibitor (NSC23766; Calbiochem, San Diego, CA) 3 h prior to infection.

Western Blot analysis of Yop injection

To analyze Yop injection by immunoblotting, cells were grown in 94 mm dishes, and then were left untreated uninfected or were infected with the E40-pBla reporter strain or the ΔYscN-pBla strain (MOI 50) as a control for 90 minutes. Cells were lysed in 200 μl PBS with 1% Triton-X (Sigma). 100 μg cell lysate protein was analyzed by SDS-PAGE and immunoblotting was performed in similar as already described for detection of Yop secretion with the exception that HRP Substrate Plus (P. J. K., Kleinbittersdorf, Germany) was used instead of ECL and in some cases film exposure was extended to 30 min for increased sensitivity.

Adhesion to cultured cells

To assay adhesion of the E40-pBla reporter strain to HeLa, GD25 and GD25-B1A cells, the respective cells were seeded on coverslips in a 24-well plate (Nunc Life Technologies, Wiesbaden, Germany). One hour after infection (MOI 50), cells were washed with PBS, fixed with 4% PFA and stained with fuchsine. All samples were prepared in duplicates, of which cells and bacteria were counted in 6 representative details.

RNA interference

For the knock-down of RhoGTPases, the predesigned and prevalidated siRNA oligonucleotides Hs Hs Rac1-6 and Hs Cdc42-7 were purchased from Qiagen (Hilden, Germany). AllStars negative control siRNA (Qiagen) was used as a nonsilencing control. 3×10⁶ HeLa cells were transfected with siRNA (5 nM) in a 12-well plate 48 h prior to infection using HiPerFect transfection reagent (Qiagen) according to the manufacturer’s fast forward protocol.

Successful knock-down was confirmed on RNA level by qPCR. Total mRNA of infected HeLa cells was extracted using RNAeasy Mini Kit (Qiagen). 1 μg of total mRNA was reverse transcribed to cDNA using QuantiTect Reverse Transcription Kit (Qiagen). Real-time qPCR was performed on a LightCycler 480 (Roche Diagnostics, Mannheim, Germany) using QuantiTect SYBR Green PCR Kit and the respective QuantiTect Primer Assays (Rac1 1, RhoA 1, Cdc42 2 and GAPDH 2). Relative expression
levels were calculated using the ΔΔCT method [65]. Expression levels of the target genes were normalized to glyceraldehyde-3-phosphate dehydrogenase RNA expression. Thus, we could demonstrate at least 75% knock-down by the used siRNAs at mRNA expression level (data not shown).

In vivo infections
C57BL/6 mice were purchased from Harlan Winkelmann (Borchen, Germany), TNF-Rp55−/− [66] and IFN-γR−/− [67] mice with a genetic C57BL/6 background were bred under specific pathogen-free conditions. Animal experiments were performed according to german law with permission of the Regierungspräsidium Tübingen. The experiments were performed with 6–8-week old female mice. To enable profound infection, mice were in parallel treated with desferrioxamine. After two days, mice were sacrificed by CO2 asphyxiation and their spleens were surgically removed and placed in ice-cold HBSS (Ca2+ and Mg2+ free Hanks’ balanced salt solution) (Biochrom) supplemented with 2% v/v fetal calf serum (FCS) (Sigma Chemical) and 10 nM HEPES buffer (Biochrom). Single cells were obtained by forcing the organs with a 5 ml syringe pestle (Falcon; BD Labware, Franklin Lakes, USA). Cell suspensions were washed twice with ice-cold HBSS. Red blood cells were lysed from spleen samples by incubating the cell suspensions for 5 min at room temperature in lysis buffer (170 mM Tris, 160 mM NH4Cl, 10 mM HEPES buffer, pH 7.4) followed by two washes in ice-cold HBSS.

Detection of Bla-activity by microscopy
For the detection of Bla activity by immunofluorescence microscopy, HeLa cells were washed once with PBS and covered with 1× CCF4-AM staining solution supplemented with probenecid, prepared according to the manufacturer’s instructions. Cells were incubated 30 min at a dark place at room temperature prior to observation with an Axiovert 200 microscope. Pictures were taken with an Axioscam HRc and Axiovision 4.4 Software was used to capture the shots and to produce overlay images (Carl Zeiss Microimaging, Esslingen, Germany). Filter sets for individual staining signal were prepared and stained by an immunoperoxidase method using Mayer’s hemalaun, mounted, and assessed microscopically on a BX51 microscope (Olympus Optical Co, Leinfelden, Germany) by two independent investigators. Pictures were taken with a DP71 camera and analysed with cell® software (Olympus). Immunostaining of controls was negative for all groups tested.

Statistics
If not otherwise stated, the means and standard deviations (SD) of data derived by cell culture experiments are calculated from four independent experiments. The number of mice which were used to calculate means and SD in mouse infection experiments is indicated in the manuscript or figure legends. Statistical analyses were performed using one-way ANOVA analyses with either Dunnett (comparison of control group with other groups) or Bonferroni corrections (comparison of all groups) as indicated in the figure legends or the text by using Graph Pad Prism software (GraphPad Software, La Jolla, USA). This software was also used to fit non-linear regression models.

Supporting Information
Figure S1 Example for the flow cytometry analysis after in vivo infection. An example of the analyses for the determination of blue cells in splenocytes after E40-pOva and E40-pBla infection of mice is shown. (A) Forward and side scatter was analyzed of all dots measured. (B) High green fluorescence (green) characterizes uptake of CCF4 and therefore viability of cells, (C) shows as green dots that the green fluorescent viable cells are mostly found in R2. Gating with R1 and R2 results in the cell populations shown in D. (D) Cells in gate R3 were defined as blue cells. Found at: doi:10.1371/journal.ppat.1000551.s001 (0.30 MB TIF)

Figure S2 Analysis of B cell subpopulations after in infection in vitro. Splenocytes were cultured and infected for one hour with or without E40-pBla (MOI 50). Cells were harvested and stained with anti-CD19-APC, anti-CD21-PE-Cy7 and then incubated with CCF4. Cells were analyzed by flow cytometry. Analysis was performed as followed. (A) Viable B cells were defined by gating using R1 & R2 & R3. All viable CD19+ cells (total) or blue CD19+ cells (R4) were analyzed for CD21 and CD23 expression. Percentages of CD21− (NFB, newly formed B cells), CD21+CD23− (MZ, marginal zone B cells) and CD21+CD23+ (FO, follicular B cells) are indicated by numbers for this experiment. In total two experiments with similar results were performed. Mean and SEM of (B) the percentages of CD19+ cells, (C) the percentages of blue cells of the indicated B cell populations and (D) the percentages of the composition of total CD19+ blue cells are shown.
**Figure S3** Analysis of activation of B cells in cell culture. Splenocytes were cultured and infected for one hour with or without E40-pBla. Cells were harvested and stained with anti-CD19-APC, anti-CD69-PE-Cy7 and MHC-II-APC-Cy7 and then incubated with CCF4. Cells were analyzed by flow cytometry. Analysis was performed in the following manner: (A) viable B cells were defined by gating using R1 & R2 & R3. (B) B cells were then further analysed for CD69 expression levels of all cells using additionally gate R3 (green+ blue+ cells), R6 (green+ blue− cells) or R7 (total). Histograms for CD69 expression of uninfected or E40-pBla infected cells are shown. Numbers indicate mean fluorescence intensities (MFI) for one representative experiment. Found at: doi:10.1371/journal.ppat.1000551.s003

**Figure S4** Infection of splenocytes of TNFR−/−, IFN-γR−/− and C57BL/6 mice in cell culture. Splenocytes were infected with indicated MOI for one hour and subsequently cells were stained with CCF4-AM and analyzed by flow cytometry. Data summarize three independent experiments.

**References**

1. Smego RA, Frean J, Koomhof HJ (1999) Yersiniosis I: microbiological and clinicopathological aspects of plague and non-plague Yersinia infections. Eur J Clin Microbiol Infect Dis 18: 1–15.
2. Koomhof HJ, Smego RA Jr, Nicol M (1999) Yersiniosis II: The pathogenesis of Yersinia infections. Eur J Clin Microbiol Infect Dis 18: 87–112.
3. Cornelis GR (2002) Yersinia type III secretion: send in the effectors. J Cell Biol 158: 401–406.
4. Hecue CJ (1998) Type III protein secretion systems in bacterial pathogens of animals and plants. Microbiol Mol Biol Rev 62: 379–433.
5. Cornelis GR (2006) The type III secretion injection. Nat Rev Microbiol 4: 811–825.
6. Galan JE, Wolfratz H (2006) Protein delivery into eukaryotic cells by type III secretion machines. Nature 444: 567–573.
7. Edqvist PJ, Aili M, Liu J, Francis MS (2007) Minimal YopB and YopD translocator secretion by Yersinia is sufficient for Yop- effector delivery into target cells. Microbes Infect 9: 224–233.
8. Hakansson S, Bergman T, Vanoosteghem JC, Cornelis G, Wolfratz H (1993) YopD and YopD constitute a novel class of Yersinia Yop proteins. Infect Immun 61: 71–80.
9. Trulsch K, Sporleder T, Igwe EI, Russmann H, Heesemann J (2004) Contribution of the major secreted yops of Yersinia enterocolitica O8 to pathogenicity in the mouse infection model. Infect Immun 72: 5237–5244.
10. Roland A, Cornelis GR (1990) Role of YopH in suppression of tumor necrosis factor alpha release by macrophages during Yersiniosis infection. Infect Immun 66: 1878–1884.
11. Mihklerjev S, Krisy G, Li Y, Wang Y, Hall HL, Goldsmith EJ, Orth K (2006) Yersinia YopH acetylates and inhibits kinase activation by blocking phosphorylation. Science 312: 1211–1214.
12. Thielis A, Wolf A, Doerrler T, Grass GA, Matsusho K, Autenrieth IB, Bohm E, Sakurai H, Niedenthal R, Resch K, Krauch M (2006) The Yersinia enterocolitica effector YopP inhibits host cell signalling by inactivating the protein kinase TAK1 in the IL-1 signalling pathway. EMBO Rep 7: 838–844.
13. Mittal R, Peak-Chew SY, McMahon HT (2006) Acetylation of MEK2 and I kappa B kinase (IKK) activation loop residues by YopH inhibits signaling. Proc Natl Acad Sci U S A 103: 18574–18579.
14. Orth K, Xu Z, Mudgett MB, Ba QZ, Palmer LE, Bliska JB, Mangel WF, Staskawicz B, Dixon JE (2000) Disruption of signaling by Yersinia effector YopJ, a ubiquitin-like protein protease. Science 290: 1594–1597.
15. Autenrieth SE, Soldanova I, Rosenmann R, Gunst D, Zahir N, Kracht M, Ruckdeschel K, Wagner H, Borgmann S, Autenrieth IB (2007) Yersinia enterocolitica YopP inhibits MAP kinase-mediated antigen uptake in dendritic cells. Cell Microbiol 9: 425–437.
16. Bohm E, Muller S, Lauber J, Gheffers R, Speer N, Georgiev P, Buer J, Zell A, Autenrieth IB (2004) Gene expression patterns of epithelial cells modulated by pathogenicity factors of Yersinia enterocolitica. Cell Microbiol 6: 129–141.
17. Erfurth SE, Grobner S, Kramer U, Gunst DS, Soldanova I, Schaller M, Autenrieth IB, Borgmann S (2004) Yersinia enterocolitica induces apoptosis and inhibits surface molecule expression and cytokine production in murine dendritic cells. Infect Immun 72: 7045–7054.
18. Denecker G, Declercq W, Geuijen CA, Boland A, Benabdillah R, van Gorp M, Sory MP, Vandenabeele P, Cornelis GR (2001) Yersinia enterocolitica YopP-induced apoptosis of macrophages involves the apoptotic signaling cascade upstream of hI. J Biol Chem 276: 19706–19714.
19. Grobner S, Autenrieth SE, Soldanova I, Gunst DS, Schaller M, Bohm E, Muller S, Leverkus M, Wesselborg S, Autenrieth IB, Borgmann S (2006) Yersinia Yop-induced apoptotic cell death in murine dendritic cells is partially independent from action of caspases and exhibits necrosis-like features. Cell Microbiol 8: 129–141.
20. Grobner S, Autenrieth SE, Soldanova I, Gunst DS, Schaller M, Bohm E, Muller S, Leverkus M, Wesselborg S, Autenrieth IB, Borgmann S (2006) Yersinia Yop-induced apoptotic cell death in murine dendritic cells is partially independent from action of caspases and exhibits necrosis-like features. Apoptosis 11: 1959–1968.
21. Viboud GI, Bliska JB (2005) Yersinia outer proteins: role in modulation of host cell signaling responses and pathogenesis. Anna Rev Microbiol 59: 69–89.
22. Gerke C, Fallow S, Chien YH (2005) The adaptor molecules LAT and SLP-76 are specifically targeted by Yersinia to inhibit T cell activation. J Exp Med 201: 561–571.
23. Alonso A, Bottini N, Breuninger S, Tschopp J, Husstedt CT, Heesemann J (2005) Yersinia outer protein P of Yersinia enterocolitica simultaneously blocks the nuclear factor-kappa B pathway and exploits lipopolysaccharide signaling to trigger apoptosis in macrophages. J Immunol 166: 1823–1831.
24. Day JB, Ferrari P, Plano GV (2003) Infection of splenocytes of TNFR−/−, IFN-γR−/− and C57BL/6 mice in cell culture. Splenocytes were infected with indicated MOI for one hour and subsequently cells were stained with CCF4-AM and analyzed by flow cytometry. Data summarize three independent experiments.
34. Schmid Y, Grassl GA, Buhler OT, Skurnik M, Autenrieth IB, Bohn E (2004) Yersinia enterocolitica adhesin A induces production of interleukin-8 in epithelial cells. Infect Immun 72: 6780–6789.

35. Mejia E, Blücker JB, Viboud GI (2000) Yersinia controls type III effector delivery into host cells by modulating Rho activity. PLoS Pathog 4: e3. doi:10.1371/journal.ppat.0040003.

36. Autenrieth IB, Beer M, Bohn E, Kaufmann SHE, Heesemann J (1994) Immune responses to Yersinia enterocolitica in susceptible BALB/c and resistant C57BL/6 mice: an essential role for gamma interferon. Infect Immun 62: 2590–2599.

37. Autenrieth IB, Heesemann J (1992) In vivo neutralization of tumor necrosis factor alpha and interferon-gamma abrogates resistance to Yersinia enterocolitica in mice. Med Microbiol Immunol 181: 333–338.

38. Bohn E, Autenrieth IB (1996) IL-12 is essential for resistance against Yersinia enterocolitica by triggering IFN-gamma production in NK cells and CD4+ T cells. J Immunol 156: 1458–1468.

39. Bohn E, Schmitt E, Riether C, Schulte R, et al. (1998) Ambiguous role of interleukin-12 in Yersinia enterocolitica infection and resistant mouse strains. Infect Immun 66: 2213–2220.

40. Di Genaro MS, Waidmann M, Kramer U, Autenrieth IB, Heesemann J (2001) Beta 1 integrin-dependent and -independent polymerization of fibronec-tin in the interferon-gamma receptor [see comments]. Science 259: 1742–1745.

41. Eitel J, Dersch P (2002) The YadA protein of Yersinia pseudotuberculosis secretes virulence factors into the extracellular milieu. Mol Microbiol 31: 1619–1629.

42. Gascon JP, Autenrieth IB, Heesemann J, Roos D (1993) Yersinia enterocolitica biogroup 1A strain for the construction of cloning vectors. J Biotechnol 79: 63–72.

43. Gascon JP, Autenrieth IB, Heesemann J, Roos D (1993) Yersinia enterocolitica biogroup 1A strain for the construction of cloning vectors. J Biotechnol 79: 63–72.

44. Gascon JP, Autenrieth IB, Heesemann J, Roos D (1993) Yersinia enterocolitica biogroup 1A strain for the construction of cloning vectors. J Biotechnol 79: 63–72.

45. Gascon JP, Autenrieth IB, Heesemann J, Roos D (1993) Yersinia enterocolitica biogroup 1A strain for the construction of cloning vectors. J Biotechnol 79: 63–72.

46. Gao Y, Dickerson JB, Guo F, Zheng J, Zheng Y (2004) Rational design and construction of urease-negative mutants of Yersinia enterocolitica serotypes O:3 and O:8: role of urease in virulence and arthritogenicity. Infect Immun 68: 942–947.

47. Gao Y, Dickerson JB, Guo F, Zheng J, Zheng Y (2004) Rational design and construction of urease-negative mutants of Yersinia enterocolitica serotypes O:3 and O:8: role of urease in virulence and arthritogenicity. Infect Immun 68: 942–947.

48. Gao Y, Dickerson JB, Guo F, Zheng J, Zheng Y (2004) Rational design and construction of urease-negative mutants of Yersinia enterocolitica serotypes O:3 and O:8: role of urease in virulence and arthritogenicity. Infect Immun 68: 942–947.

49. Gripenberg-Lerche C, Zhang L, Ahtonen P, Toivanen P, Skurnik M (2000) Yersinia enterocolitica: role of the Yersinia virulence plasmid and modulation by the iron-chelator desferrioxamine B. J Infect Dis 170: 140–150.

50. Gripenberg-Lerche C, Zhang L, Ahtonen P, Toivanen P, Skurnik M (2000) Yersinia enterocolitica: role of the Yersinia virulence plasmid and modulation by the iron-chelator desferrioxamine B. J Infect Dis 170: 140–150.

51. Ekdahl J, Hesseman J, Röjdler H, Autenrieth IB (1994) Interaction of polymorphonuclear leukocytes with Yersinia enterocolitica: role of the Yersinia virulence plasmid and modulation by the iron-chelator desferrioxamine B. J Infect Dis 170: 140–150.

52. Ekdahl J, Hesseman J, Röjdler H, Autenrieth IB (1994) Interaction of polymorphonuclear leukocytes with Yersinia enterocolitica: role of the Yersinia virulence plasmid and modulation by the iron-chelator desferrioxamine B. J Infect Dis 170: 140–150.

53. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

54. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

55. Heesemann J (1987) Chromosomal-encoded siderophores are required for mouse virulence of enteropathogenic Yersinia species. FEMS Microbiol Lett 48: 229–233.

56. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

57. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

58. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

59. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

60. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

61. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

62. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

63. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

64. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.

65. Heesemann J, Gross U, Schmidt N, Laufs R (1986) Immunochemical analysis of macrophages. J Infect Dis 172: 490–496.