Reduced acute myocardial ischemia–reperfusion injury in IL-6-deficient mice employing a closed-chest model

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Abstract

Objective and design We examined the role of IL-6 in the temporal development of cardiac ischemia–reperfusion injury employing a closed-chest I/R model.

Materials/methods Infarction, local and systemic inflammation, neutrophil infiltration, coagulation and ST elevation/resolution were compared between wild-type (WT) and IL-6-deficient (IL-6−/−) mice after 1 h ischemia and 0, 1/4, 3, and 24 h reperfusion.

Results IL-6 deficiency reduced infarct size at 3 h reperfusion (28.8 ± 4.5 % WT vs 17.6 ± 2.5 % IL-6−/−), which reduction persisted and remained similar at 24 h reperfusion. Serum Amyloid A was reduced at 24 h reperfusion only (57.5 ± 4.9 WT vs 24.8 ± 5.6 ug/ml IL-6−/− mice). Cardiac cytokines (IL-6, IL-1β and TNFα) peaked at 3 h reperfusion, but IL-1β and TNFα levels were unaffected by IL-6 deficiency. Significant neutrophil influx was only detected at 24 h reperfusion and was similar for WT and IL-6−/−. Tissue factor peaked at 24 h reperfusion, whereas fibrin/fibrinogen peaked at 3 h reperfusion and was completely resolved at 24 h reperfusion; both coagulation factors were unaltered by IL-6 deficiency. Prolonged ST elevation was observed during ischemia that completely resolved for both genotypes at early reperfusion.

Conclusions The data suggest that, in the absence of major surgical intervention, IL-6 contributes to the development of infarct size in the early phase of reperfusion; this contribution did not depend on neutrophil influx, IL-1β and TNFα, tissue factor and fibrin.

Keywords Inflammation · IL-6 · Closed-chest model · Ischemia/reperfusion injury · Heart

Introduction

Acute myocardial ischemia–reperfusion (I/R) is associated with an inflammatory response that may contribute to I/R-induced injury of the heart [1, 2]. One important pleiotropic inflammatory mediator that is released quickly upon cardiac I/R is interleukin (IL)-6 [3, 4]. Clinical studies have shown that chronically elevated IL-6 levels in blood are independently associated with acute coronary syndromes, suggesting an etiologic role for this cytokine in the process of myocardial injury [5, 6]. However, several pre-clinical animal studies have refuted IL-6 as major contributor to
We, therefore, measured infarct sizes, tested pronounced pro-inflammatory and pro-coagulant effect. Smaller infarct sizes in IL-6 deficient mice due to a less activation of the coagulation system. We expected to find does so through alteration in inflammatory pathways and acute myocardial ischemia–reperfusion injury, and (2) it type and IL-6 after an acute episode of I/R were similar between wild-type and IL-6−/− mice [9–11]. Recently, Samanta et al. have shown that IL-6 did contribute to adverse remodeling at 35 days following I/R [8]. However, these studies performed I/R immediately following the surgical preparative intervention of chest opening and puncturing of the heart to allow the ligation of the LAD. This surgical intervention significantly raises the baseline level of cytokines where-upon the effects of I/R-induced cytokine production are studied [12]. In a previous study we demonstrated that 75 % of the inflammatory cytokine response to acute I/R in such an open-chest model could be ascribed to the chest opening. In addition, the opening of the chest was associated with priming of the NLRP3 inflammasome, which priming was not observable in the closed-chest model [13]. Knowing that effects of inflammatory mediators are highly dependent on the actual concentration of the cytokine and the state of the innate immune system (e.g., [1]), it is, therefore, still unknown whether IL-6 affects acute cardiac injury in the absence of major surgery, i.e., in a closed-chest model of cardiac I/R. Such a closed-chest model better reflects the condition of acute cardiac syndromes as occurring in AMI patients and/or during PCI procedures, whereas the open-chest model better mirrors cardiac surgical procedures. Therefore, in the present study we examined to what extent IL-6 affects acute myocardial I/R injury in a closed-chest mouse model.

IL-6 belongs with tumor necrosis factor (TNF)α and IL-1β to the pro-inflammatory cytokines and may, therefore, contribute to increases in serum amyloid A (SAA) [14] in the systematic circulation and to neutrophil influx into the injured cardiac tissue [15]. In addition, it has been suggested that IL-6 contributes to activation of the coagulation system in experimental models of endotoxemia, due to the existence of cross-talk between inflammation and coagulation [16, 17]. Such an increased coagulation, when associated with microvascular thrombosis, is then a possible mechanism through which IL-6 can increase infarct size after myocardial ischemia–reperfusion [18]. Possible changes in coagulation may then impact the resolution of ischemia during reperfusion due to changes in no-flow regions within the heart, as reflected by a divergent ST-segment resolution [19, 20].

In this study we hypothesized that (1) IL-6 contributes to acute myocardial ischemia–reperfusion injury, and (2) it does so through alteration in inflammatory pathways and activation of the coagulation system. We expected to find smaller infarct sizes in IL-6 deficient mice due to a less pronounced pro-inflammatory and pro-coagulant effect. We, therefore, measured infarct sizes, tested pro-inflammatory cardiac cytokine levels and systemic SAA levels, used immunohistochemistry to evaluate neutrophil influx and activation of the coagulation system (tissue factor and fibrin/fibrinogen), and used ECG analysis as a surrogate marker of no-flow in a closed-chest mouse model [21] after 1 h ischemia and ½, 3 or 24 h reperfusion.

Materials and methods

Animals

IL-6−/− mice were originally generated by Kopf et al. [22]. Eight- to 10-week-old male IL-6−/− mice (backcrossed onto a C57BL/6 background for at least 6 generations) and inbred C57BL/6 mice (for controls) (Jackson Laboratory; Bar Harbor, ME) were used in this study. They were housed in open topped cages at constant temperature (21–22 °C), humidity (50 ± 10 %) and light cycle (12 h dark–light). Rodent chow (AM-II 10 mm, Hope Farms, Woerden, The Netherlands) and drinking water were available ad libitum. Experiments were approved under number DIX 228 by the animal ethics committee of the Academic Medical Center, Amsterdam, The Netherlands, and comply with US National Institutes of Health or European Commission guidelines. The flowchart (Fig. 1) shows the number of mice in each analysis.

Preparative surgery for I/R in closed chest

Stable anesthesia (with stable hemodynamic parameters) was induced by 10 ml FFM/kg BW i.p. (FFM: Fentanyl-Fluanisone-Midazolam) [23, 24]. Hypnorm® (VetaPharma, Leeds, UK) contains 0.315 mg Fentanyl citrate/ml and 10 mg Fluanisone/ml. Midazolam® (Actavis, Iceland) contains 5 mg Dormicum/ml. Both Hypnorm and Midazolam were 1:1 diluted in H2O prior to their 1:1 mixing. Anesthesia was maintained by bolus injections of ½ times the initial FFM volume. Paw withdrawal reflex and tail pinch were used to monitor the adequacy of anesthesia. After an induction time of 5 min, a preparative surgery was started as described previously [21]. Briefly, the skin of the throat was cut to visualize the trachea. The mice were intubated with an Intramedic PE-90 tube (Becton Dickinson, Sparks, Md), and volume controlled ventilated (CIV101 ventilator, Columbus Instruments, Ohio) with room air supplied with 40 % O2. Body temperature was maintained between 36.6 and 37.4 °C. Lateral thoracotomy was performed between the third and fourth rib, while the rectus abdominis and pectoralis were kept intact and put aside. The pericardium was locally opened, and a U-bend 8–0 prolene suture was put under the LAD approximately 2 mm lower than the tip of the left auricle,
similarly as reported by Michael et al. [2] and Xu et al. [25]. The endings were tunneled in parallel through a 2 mm Intramedic PE-10 tube (Becton Dickinson, Sparks, Md) and separately brought outside between the adjacent ribs. The chest was closed under application of gentle pressure on both sides of the thorax to expel air, and the endings of the 8–0 prolene suture were hidden s.c. Post-operative analgesia was induced by 0.1 mg buprenorphine/kg s.c. (Temgesic/C210, Schering-Plough, Maarssen, The Netherlands). Sterile 0.5 ml 0.9 % saline i.p. was administered as fluid. Mice were allowed to recover in a temperature-controlled room of 30°C for 1 day and at room temperature for >8 days.

**Ischemia–reperfusion**

9–11 days after the preparative surgery, when the inflammatory reaction had subsided [12, 21] the mice were reanesthetized with FFM, and allowed to freely breathe under 40 % oxygen-supplemented air. ECG (Einthoven lead I) was recorded by AgCL coated-silver needles. Output signals were amplified by a custom-made amplifier, sampled at 1 kHz, and band pass filtered at 100, 50 Hz and DC, and directly transferred to LabView5.0 (National Instruments, Austin, TX, USA). The skin was opened over the old scar, and the LAD (8–0 prolene) sutures were tightened for 1 h to achieve ischemia as verified by ECG recording. Release of the sutures, again confirmed by ECG, resulted in reperfusion. Mice were killed at different time points after 0, ½, 3, or 24 h reperfusion. Sham-treated mice had identical surgical procedures as the experimental group except that the sutures were not pulled. Mice in the 24 h reperfusion group were administered 0.1 mg buprenorphine/kg s.c. (Temgesic®, Schering-Plough, Maarssen, The Netherlands) after 30 min reperfusion and housed in a temperature-controlled room of 30°C. All mice were killed after reperfusion under FFM anesthesia.

**Blood sampling**

After each experiment, the mice were heparinized via the tail vein (200 Units heparin (Leo Pharma, Breda, Netherlands) in 0.2 ml sterile saline) to prevent coagulation in the tissues [26]. Blood was collected from the vena cava into a plastic syringe containing 0.1 volume of 3.2 % sodium citrate. After centrifugation plasma samples were stored at −80°C until further analysis.

**Infarct size**

To identify the area at risk (AAR) and area of infarction (AOI) within the left ventricle of WT and IL-6−/− mice after 3 and 24 h reperfusion, hearts were subjected to Evans blue and 2,3,5-triphenyl-tetrazolium chloride (TTC) staining [2]. Briefly, after exsanguination the chest was opened via medial section, on the right side of the septum. The sutures were tied to definitely occlude the LAD. A 23G1 Gauge blunted needle (Becton–Dickinson, Etten-Leur, The Netherlands) attached to a 2-ml syringe filled with 4°C cardioplegic solution was cannulated through the aorta into the LV lumen and attached to the aorta. The heart was excised from the body and (retrogradely) infused with cardioplegia to rinse. Next, the needle was attached to a 1-ml syringe and the heart was infused with 50–100 µl 2 % Evans Blue (Sigma, St. Louis, MO, USA) to negatively mark the area at risk (AAR). The heart was cooled for several min at −20°C after infusion of type VII low gelling temperature agarose (Sigma, St. Louis, MO, USA). The needle was removed and the heart was serially cut into transverse sections. The first section contained the suture around the LAD. To differentiate between the viable and necrotic areas of the myocardium within the AAR, the three 0.8 mm sections, the apex, and the base of the heart were soaked between Whatman filter paper and stained by 1.5 % TTC (Sigma-Aldrich, Steinheim, Germany) in

![Fig. 1 Flowchart with time frame, numbers of mice used and analyses performed](image-url)
phases after 0, /C145 systems, Abingdon, UK). Formed using commercially available ELISA kits (R&D level 0.50; 0.63; and 0.60 ng/mg, respectively) were per-

After 24 h reperfusion, IL-6, IL-1β, and TNFα levels were in duplicate determined in plasma and in supernatants from control hearts, sham hearts, and from ischemic and non-ischemic parts (hearts were longitudinally split into two parts that contained the ischemic area and the area not at risk to distinguish the local inflammatory responses). Measurements of IL-6, IL-1β, and TNFα levels (detection level 0.50; 0.63; and 0.60 ng/mg, respectively) were performed using commercially available ELISA kits (R&D systems, Abingdon, UK).

SAA levels were singularly determined in remaining plasmas after 0, ½, 3, and 24 h reperfusion (detection level 55 ng/ml) (Biosource International, Inc. Camarillo, CA, USA).

Cytokines (in hearts and plasma), and SAA (in plasma)

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Homogenate preparation

After excsanguinations, hearts were homogenized (1 vol) in Greenburger lysis buffer (300 mM NaCl, 15 mM Tris, 2 mM MgCl2, 2 mM Triton(X-100), Pepstatin A, Leupeptin, Aprotinin (20 ng/ml), pH 7.4) (9 vol). After centrifugation, the supernatant was frozen at −80 °C for cytokine measurements.

Image analysis

The hematoxylin and eosin section with the largest visible I/R damage was the starting point to select 3 adjacent sections a heart for immunohistochemistry; The number of Ly-6G-positive cells, and the expression of tissue factor and fibrin/fibrinogen was investigated in the same areas and quantified by digital image analysis. In short, stained sections were scanned using the Philips IntelliSite Ultra Fast Scanner (Philips Digital Pathology Solutions, Best, The Netherlands). The mid-left ventricular free wall (0.33–1.2 mm [2]) was selected (endocardium and peri-

Endogenous peroxidase activity was blocked by 1.5 % H2O2 in PBS for 30 min at room temperature, and washed in PBS. Non-specific binding sites were blocked with TENG-T [10 mM Tris, 5 mM EDTA, 0.15 M NaCl, 0.25 % gelatin, 0.05 % (vol/vol) Tween-20, pH 8.0] or 10 % normal goat serum (Dako, Glostrup, Denmark) for 30 min at RT, and then washed in PBS. Sections were incubated at 4 °C overnight with the different primary antibodies. The following antibodies were used: Ly6 for neutrophils (Pharmingen, San Diego, CA, USA), Tissue Factor (1:250 anti-tissue factor) and fibrin (Biotinylated goat anti-mouse fibrinogen (Accurate Chemical & Scientific Corporation, Westbury, NY) recognized fibrin/fibrinogen deposition). In each case, controls were treated with isotypic control IgG. Sections stained for Ly6G and fibrin were subsequently incubated with HRP conjugated swine anti rabbit immunoglobulins and enzyme activity was detected using 3,3′-diaminobenzidine-tetra-hydrochloride (DAB, Sigma-Aldrich, St. Louis, MO) or 3-amino-9-ethylcarbazole (AEC, Sigma-Aldrich) as a sub-

liminal fibrin/fibrinogen) and analyzed.

Tissue sampling for histology and immunohistochemistry

After ½, 3, and 24 h reperfusion, WT mice and IL-6−/− were killed by excsanguinations under anesthesia (see paragraph “Blood sampling”). Hearts were taken from the chest, and transversally cut at the side where the suture was loosely saved after induction of reperfusion. The apex was removed and the mid-section was fixed in 4 % paraformaldehyde for 6 h at room temperature, and embedded in paraffin using automatic embedding equipment (Tissue Tek, Miles Scientific Inc., USA). 4 μm sections were obtained and every 30th slice was stained with hematoxylin and eosin, which resulted in 10–13 stained slices a heart. For immunohistochemistry, the tissue sections were deparaffinized, rehydrated, and washed with water and PBS. Endogenous peroxidase activity was blocked by 1.5 % H2O2 in PBS for 30 min at room temperature, and washed in PBS. Non-specific binding sites were blocked with TENG-T [10 mM Tris, 5 mM EDTA, 0.15 M NaCl, 0.25 % gelatin, 0.05 % (vol/vol) Tween-20, pH 8.0] or 10 % normal goat serum (Dako, Glostrup, Denmark) for 30 min at RT, and then washed in PBS. Sections were incubated at 4 °C overnight with the different primary antibodies. The following antibodies were used: Ly6 for neutrophils (Pharmingen, San Diego, CA, USA), Tissue Factor (1:250 anti-tissue factor) and fibrin (Biotinylated goat anti-mouse fibrinogen (Accurate Chemical & Scientific Corporation, Westbury, NY) recognized fibrin/fibrinogen deposition). In each case, controls were treated with isotypic control IgG. Sections stained for Ly6G and fibrin were subsequently incubated with HRP conjugated swine anti rabbit immunoglobulins and enzyme activity was detected using 3,3′-diaminobenzidine-tetra-hydrochloride (DAB, Sigma-Aldrich, St. Louis, MO) or 3-amino-9-ethylcarbazole (AEC, Sigma-Aldrich) as a sub-

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Cardiac microscopy and immunohistochemistry

For immunohistochemistry, the tissue sections were deparaffinized, rehydrated, and washed with water and PBS. Endogenous peroxidase activity was blocked by 1.5 % H2O2 in PBS for 30 min at room temperature, and washed in PBS. Non-specific binding sites were blocked with TENG-T [10 mM Tris, 5 mM EDTA, 0.15 M NaCl, 0.25 % gelatin, 0.05 % (vol/vol) Tween-20, pH 8.0] or 10 % normal goat serum (Dako, Glostrup, Denmark) for 30 min at RT, and then washed in PBS. Sections were incubated at 4 °C overnight with the different primary antibodies. The following antibodies were used: Ly6 for neutrophils (Pharmingen, San Diego, CA, USA), Tissue Factor (1:250 anti-tissue factor) and fibrin (Biotinylated goat anti-mouse fibrinogen (Accurate Chemical & Scientific Corporation, Westbury, NY) recognized fibrin/fibrinogen deposition). In each case, controls were treated with isotypic control IgG. Sections stained for Ly6G and fibrin were subsequently incubated with HRP conjugated swine anti rabbit immunoglobulins and enzyme activity was detected using 3,3′-diaminobenzidine-tetra-hydrochloride (DAB, Sigma-Aldrich, St. Louis, MO) or 3-amino-9-ethylcarbazole (AEC, Sigma-Aldrich) as a sub-

Ly6G-positive cells were manually counted using the ImageJ cell counter and expressed as number of neutrophils/mm² [2]. Tissue factor and fibrin images were converted to 8-bit grayscale images and the immunopositivity was measured by setting an intensity threshold. The immunopositive area was expressed as a percentage of the total area.
mouse and two IL-6−/− mice seemed to have disturbing electrical noise and were excluded from the data. The resulting ECG readings of 19 WT and 18 IL-6−/− mice were averaged over 10 s recordings using software written in the Matlab (the Mathworks, USA) script language. Time alignment within a trace was performed using cross correlation with a template derived from a QRS complex. Because the morphology of the QRS complex changed as a result of the intervention it was not possible to align the differently shaped QRS complexes. Alignment of different traces was, therefore, done using the P-wave. Afterwards, the QRS complexes were aligned by hand. The amplitude of the ST-segment above the iso-electric line was measured 20 ms after the start of the QRS complex.

Statistical analysis

All measurements are reported as mean ± SE. All data (except AOI/AAR, AAR/LV, and AOI/LV) were not normally distributed, and therefore we opted for non-parametric testing using the one-way analysis of Kruskal–Wallis on medians. A P value of less than 0.05 was considered statistically significant. Statistics were performed using the Prism software package (GraphPad, San Diego, CA).

Results

Infarct sizes as demarcated by TTC and Evans blue staining showed significantly smaller infarcts in IL-6−/− mice when compared to WT mice after 3 h reperfusion: Fig. 2a shows infarct sizes of 28.8 ± 4.5 % in WT mice vs 17.6 ± 2.5 % in IL-6−/− mice after 3 h reperfusion. No further change in infarct size with IL-6 ablation was observed at 24 h reperfusion, 25.1 ± 3.0 % in WT mice vs 14.6 ± 4.4 % in IL-6−/− mice, indicating that IL-6-mediated effect on infarct development is effective during the first hours of reperfusion. The difference in infarct size between genotypes cannot be ascribed to differences in the area at risk (Fig. 2b). Figure 2c–f shows representative examples of TTC and Evans blue stainings.

To examine whether inflammation was constricted to the ischemic part of the heart only, we first compared the non-ischemic with the ischemic part of the heart for the ½ and 24 h reperfusion time points only. After ½ h reperfusion, cardiac IL-6 levels were already significantly elevated in the ischemic part of the heart as compared to the non-ischemic part in WT mice (5.6 ± 1.0 vs 1.7 ± 0.2 µg/mg; n = 6, respectively), indicating IL-6 to be a fast responder to cardiac ischemia (data not shown). After 24 h reperfusion cardiac IL-6 was only mildly elevated in the ischemic part, showing the transient behavior of IL-6 during I/R (ischemic 3.0 ± 0.2 vs non-ischemic 1.5 ± 0.1 µg/mg; n = 8) (data not shown). Thus, the lack of increase in IL-6 in the non-ischemic part of the heart throughout the experiment indicates that I/R-induced inflammation remains confined to the ischemic part of the heart. Figure 3 demonstrates levels of cardiac cytokines in the ischemic part of the heart for WT and IL-6−/− mice at different times of reperfusion. IL-6 levels peaked at 3 h reperfusion, and were undetectable in IL-6−/− hearts (Fig. 3a). A similar time pattern was observed for IL-1β with a peak at 3 h reperfusion (Fig. 3b). No differences in IL-1β were observed between WT and IL-6−/− at any time point, suggesting that IL-6 is without effect on IL-1β levels. Cardiac TNFα also peaked at 3 h reperfusion, with no differences in TNFα levels between WT and IL-6−/− mice at any time of reperfusion (Fig. 3c). Cardiac IL-6, IL-1β and TNFα levels in ischemic area and the non-ischemic area were not different in sham groups at 0, ½ and 24 h reperfusion (data not shown), indicating that the temporal

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changes in cytokines were due to the I/R intervention per se.

Next we examined whether the omission of IL-6 affected the acute phase protein response, by monitoring the production of SAA (produced by liver). Plasma SAA levels after 0 and 1/2 h reperfusion in WT (n = 4 and 12) and IL-6−/− (n = 6 and 12) mice were undetectable. SAA levels increased significantly in WT mice after 3 h and 24 h reperfusion and were highest after 24 h reperfusion (Fig. 4). SAA was also increased at 3 h reperfusion in IL-6−/− mice, however, in contrast to WT mice, no further increase in SAA was observed at 24 h reperfusion; resulting in significantly decreased SAA in IL-6−/− mice at 24 h reperfusion. These data indicate that the production of the acute phase response protein SAA is partly, but not solely, dictated by the rise in IL-6.

Having established that IL-6 affected the acute development of cardiac infarction and the rise in the systemic inflammatory marker SAA, we now examined whether IL-6 plays a role in neutrophil influx and coagulation activation following cardiac I/R through immunohistochemical analysis (Fig. 5). Typical examples of staining for neutrophil, tissue factor and fibrin/fibrinogen for WT and IL-6−/− are demonstrated in Fig. 5a–f. The influx of neutrophils in our model of closed-chest cardiac I/R shows a later response as compared to the cytokine response. Influx of neutrophils at 1/2 h was non-detectable, and only very minimal present at 3 h reperfusion. At 24 h reperfusion there was a significantly increased influx of neutrophils, however, this influx was similar between WT and IL-6−/− mice (Fig. 5g). Tissue factor was already detectable at 1/2 h of reperfusion and demonstrated a non-significant increase with prolongation of reperfusion time, without any significant effect of IL-6 on this development of tissue factor during reperfusion (Fig. 5h). Finally, immunostaining for fibrin/fibrinogen also revealed the early presence of this coagulation factor that peaked at 3 h reperfusion, but completely disappeared at 24 h reperfusion, with no differences between WT and IL-6−/− (Fig. 5i). Data suggest that the cytokine IL-6 is not involved in the early
recruitment of neutrophils or the activation of the coagulation system in the early reperfusion phase when employing a closed-chest model of cardiac I/R.

Finally, we applied ECG recording during cardiac I/R to monitor ST elevation as surrogate marker of degree of ischemia and successful and complete reperfusion. Figure 6 shows a summary of the averaged ST-segment elevation during ischemia, and its resolution during the early reperfusion phase for WT (Fig. 6a) and IL-6/−/− (Fig. 6b) mice. Pulling of the sutures induced an immediate ST-segment elevation that peaked after 30 s. The ST-segment elevation showed a short lasting normalization at 2 min ischemia and increased for a second time to reach its maximum after 20 min ischemia. The elevation persisted during the remaining ischemic period. Releasing of the sutures induced a short episode of partial ST-segment resolution that lasted approximately 30 s. Next the ST-segments were again elevated at 1½ min at a level that was higher than during ischemia. This temporal increase disappeared within 5 min and the ST-segment resolution was complete after 30 min for both WT and IL-6/−/− mice. In none of the mice a secondary rise or re-elevation in ST-segments was observed during reperfusion.

Fig. 5 Immunohistochemical analysis of the left ventricular wall of WT (a–c) and IL-6/−/− (d–f) mice after I/R injury. a, d Neutrophils (Ly6G) after 24 h reperfusion; b, e tissue factor after 24 h reperfusion; c, f fibrin/fibrinogen after 3 h reperfusion. Bar 100 μm.

Quantitative analysis of neutrophils (g), tissue factor (h) and fibrin/fibrinogen (i) after ½, 3, and 24 h reperfusion. Values are mean ± SEM; black dots represent WT mice; open dots represent IL-6/−/− mice.

Fig. 6 Averaged ST-segment (mV) during 60 min ischemia (uninterrupted line) and during the early reperfusion phase (interrupted line) for a WT mice (n = 19) and b IL-6/−/− mice (n = 18)
Discussion

In this study we have demonstrated that IL-6 is one of the mediators that contributes to the development of cardiac infarct size during the early period (<1 day) of reperfusion, following an injurious period of cardiac ischemia in the absence of surgical stress. In addition, our data suggest that the observed IL-6 effects on acute cardiac infarction are not mediated through changes in IL-1β, TNFα, neutrophil influx or an altered activation of the tissue factor-thrombin pathway.

IL-6 deficiency and cardiac infarct: closed-chest versus open-chest cardiac I/R models

In our closed-chest model, the absence of IL-6 resulted in smaller infarcts. These results are in contrast to previous studies employing the open-chest model of cardiac I/R, reporting no effect of IL-6 deficiency on infarct size [9, 10]. Previous studies have demonstrated [12, 13] that the major difference between these models is the enhanced activation of inflammatory pathways; i.e., the surgical stress of chest opening resulted in inflammatory priming. We demonstrated that >75% of inflammation in the open-chest model is due to surgery, and not to the actual cardiac I/R [13]. As such, the open-chest model mimics the clinical condition of cardiac I/R during CABG procedures, whereas the closed-chest model mimics the clinical condition of cardiac I/R during PCI or thrombolytic procedures. Thus, these data indicate that under conditions of an already activated inflammatory background (due to e.g., surgery in the open-chest model, sepsis, etc.), the role of a single acute inflammatory responder such as IL-6 on acute reperfusion injury is not detectable. Although open-chest studies suggest that inflammatory pathways/mediators other than IL-6 become more critical for modulating cardiac I/R injury in these conditions (e.g., the Nlrp3 inflammasome [13], TLR4 [27, 28]), further studies are necessary to decipher the precise molecular mechanisms explaining the different outcome between the open-chest and closed-chest models. Infarct sizes in closed-chest models are also usually much smaller (5–30% of risk area; present study and Durgan et al. [29]) than infarct sizes in open-chest studies (35–60% of risk area), despite the fact that in these closed-chest models a longer duration of ischemia (45–60 min) is used as compared to the open-chest models (30 min ischemia). The decreased inflammation in the closed-chest models is thus associated with a smaller infarct size as compared to open-chest models, supporting the concept that inflammation contributes to the acute infarct development upon reperfusion.

Theoretically, the inflammatory response caused by the preparative surgery could have a preconditioning effect. However, since such preconditioning has an early window (2–3 h) and a delayed window (which begins 12–24 h later and lasts 3–4 days) of protection [30] that is much shorter than the 9–11 days pause we had chosen, it is unlikely that preconditioning could have affected our I/R results in the closed-chest model.

Inflammatory mediators and I/R-induced acute cardiac infarction

Inflammation plays a modulating role in cardiac ischemia–reperfusion, both in the early reperfusion phase during infarct development and in the post-infarct phase during cardiac remodeling [31, 32]. Early responder cytokines such as TNFα and IL-6, localized in resident mast cells and endothelial cells, are quickly released and increased in expression upon the early moments of cardiac I/R. However, their role on modulating infarct size is likely critically dependent on the precise pathophysiological conditions present. TNFα knockout mice are reported to have reduced or unaltered infarct size in the isolated heart or open-chest model [32]. IL-6−/− mice are now also reported to have reduced (present study) or unaltered infarct size [9, 10]. Deletion of IL-6 was also reported to protect against acute kidney I/R [33]. It is becoming increasingly clear that these cytokines are all part of a larger network of interacting cytokines that depending on context (e.g., background inflammation, timing in I/R processes) can activate or inhibit cell death/survival signaling cascades. One, often ignored, important context characteristic, concerns the duration of the ischemic episode; we previously demonstrated this for the cardioprotective effects of folic acid: folic acid protected against 25 min ischemia, but not against 35 min ischemia [34]. Similarly, it was shown that deletion of the macrophage migration inhibitory factor protected hearts from 60 min ischemia, but trended to increase I/R injury with much shorter ischemia [35]. It is, therefore, also possible that the protection we now describe with IL-6 deletion following 60 min ischemia is still commensurate with findings that report no effect of IL-6 deletion following only 30 min ischemia [9, 10]. Literature indicates that mechanisms contributing to I/R injury differ depending on the ischemic duration, such that the efficacy of factors affecting specific mechanisms also depends on ischemia duration.

For TNFα it has been reported that this cytokine can indeed directly contribute to cell death [36], however, no such data can be found for IL-6. It has been reported that IL-6 induces mitochondrial dysfunction (decreased ATP production, increased ROS production) in
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adipocytes [37] but, although mitochondria are crucial mediators of cardiac I/R injury, it is unknown whether similar IL-6 effects are obtained in myocardial tissue. We did find decreased amounts of the acute phase response protein SAA in IL-6<sup>−/−</sup> mice. It has been shown that SAA can function as an agonist for the innate immune receptors TLR2/4 [38, 39], which receptors are part of the signaling cascades that contribute to acute cardiac infarction [27, 28, 40]. Thus, it is possible that IL-6 contributes to infarct development indirectly through activation of TLR cellular pathways.

**IL-6 effects on infarct cannot be explained by neutrophils and coagulation**

In an attempt to further examine factors that contribute to myocardial injury in WT and IL-6<sup>−/−</sup> mice, we examined neutrophil influx by histology. It is known from literature that the induction and release of IL-6 induces the expression of intercellular adhesion molecule-1 (ICAM-1) on the myocyte surface in the viable border zone of the infarct [41]. The etiologic role of released IL-6 in stimulation of myocyte ICAM-1 was previously demonstrated by others [42], and binding of neutrophils to ICAM-1 is considered to be a causal factor in myocyte injury after ischemia [15, 43, 44]. Although we did not quantify ICAM-1 expression ourselves, neutrophils were uniformly distributed throughout the previous ischemic area both in WT and IL-6<sup>−/−</sup> mice; the influx in time and the distribution were equal in WT and IL-6<sup>−/−</sup> mice suggesting that IL-6 is not important for the early recruitment of neutrophils to reperfused ischemic tissue. This can be taken as evidence that the causal pathway IL-6-ICAM-1-neutrophil is not operational in the present model; alternatively, if operational in WT mice, it apparently can be bypassed easily by other cytokines in IL-6<sup>−/−</sup> mice. In fact, TNFα was shown previously to induce ICAM-1 and subsequent neutrophil influx instead of IL-6 or IL-1β [45]. Since TNFα levels were similar between WT and IL-6<sup>−/−</sup> mice within the first 24 h of reperfusion in our study, this could well explain the fact that neutrophil influxes appeared to be independent of IL-6. Most importantly, the similar influx of neutrophils between WT and IL-6<sup>−/−</sup> mice negates that neutrophils can explain the reduced infarction observed in the IL-6<sup>−/−</sup> mice. Moreover, the difference in the time course between infarction (complete at 3 h reperfusion, no further increase with 24 h reperfusion) and influx of neutrophils (hardly detectable at 3 h reperfusion, only at 24 h reperfusion) also makes it rather unlikely that neutrophils contributed to cardiac infarction in our model. This is commensurate with current thinking that I/R-induced infarction mostly takes place during the first 30 min reperfusion, and is dictated by mechanisms such as opening of the mitochondrial transition pore, activation of calpains and hypercontractility [46, 47].

Another element in myocardial injury that may be affected by different IL-6 levels relates to coagulation activation. IL-6 is known to be an important agonist of monocytic tissue factor induction in humans and primates, either directly, or after endotoxin stimulation [16, 17], thus we expected a role of IL-6 in inducing myocardial tissue factor and subsequent fibrin formation. In addition, previous studies using an antibody against tissue factor, active protein C or active site inhibited factor VIIa demonstrated reduction in infarcted area, thus demonstrating a possible causal relationship between coagulation and ischemic myocardial injury [18, 48, 49]. However, it has recently been suggested that these anticoagulants may also directly activate cytoprotective signaling independent of coagulation inhibition [50, 51].

We were unable to detect any differences in the presence of tissue factor or fibrin at any time of reperfusion between WT and IL-6<sup>−/−</sup> mice. A possible reason as to why IL-6 does not seem to affect coagulation parameters may be related to the rather low and short-term activation of coagulation in our closed-chest cardiac I/R model. Although there is a trend in increased tissue factor at 24 h of reperfusion, the fibrin data indicate complete disappearance of fibrin at 24 h reperfusion. The fast disappearance of fibrin suggests resolution of coagulation at 24 h of reperfusion. This is also indirectly supported by our ECG monitoring. The ischemia-induced ST elevation was quickly and equally resolved during reperfusion in both WT and IL-6<sup>−/−</sup> mice. It is, therefore, likely that IL-6 exerts its detrimental effects on acute infarct development through mechanisms other than coagulation activation or neutrophil action.

In conclusion, our data indicate that IL-6 contributes to the development of infarction, in a closed-chest model of cardiac I/R that is associated with attenuated inflammation as compared to open-chest models. The beneficial effects of IL-6 deletion on infarction cannot be explained by modification of other inflammatory mediators, neutrophil influx or coagulation activation. The effects of IL-6 are apparent within the first 3 h of reperfusion, suggesting a more direct, cellular signaling effect of IL-6 on the cellular mechanism that causes I/R-induced cell death. The present findings merit a further investigation into these direct molecular mechanisms of IL-6 related to cardiac I/R injury.

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Compliance with ethical standards

Conflict of interest  The authors declare that they have no conflict of interest.

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