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Up-regulation of Grb2-associated binder 1 promotes hepatocyte growth factor induced endothelial progenitor cell proliferation and migration

Qing Fan 1,2, Liyu Zhang 1,2, Wenjie Zhu 1,2, Sheng Xue 3, Yisheng Song 1,2, Qing Chang Corresp. 1,2

1 Qingdao University, Qingdao, China
2 Cardiovascular Surgery Department, The Affiliated Hospital of Qingdao University, Qingdao, China
3 Institute for Translational Medicine, Qingdao University, Qingdao, China

Corresponding Author: Qing Chang
Email address: changqing20671@163.com

Objectives. Grb2-associated binder 1 (Gab1), a scaffolding adaptor protein, plays an important role in transmitting key signals that control cell growth, migration, and function from multiple tyrosine kinase receptors. This study was designed to investigate the influence of upregulation of Gab1 in endothelial progenitor cells (EPCs) stimulated with hepatocyte growth factor (HGF), and the underlying molecular mechanisms.

Materials and Methods. EPCs isolated from human umbilical cord blood were identified and divided into four groups. EPCs in the Control group were cultured normally; those in the Control+HGF group were treated with HGF stimulation; those in the AD-Gab1 group were transfected with adenovirus containing the Gab1 gene but not treated with HGF stimulation; and, those in the AD-Gab1+HGF group were treated with both HGF stimulation and transfection with adenovirus containing the Gab1 gene. Subsequently, Gab1 expression and proliferation and migration ability were compared for EPCs grown under different conditions. Furthermore, we measured phosphorylation levels of three key proteins Gab1, SHP2 and ERK1/2.

Results. The AD-Gab1+HGF group had the highest expression of Gab1 and higher proliferation and migration than the other three groups.

Conclusions. Upregulation of Gab1 promoted HGF-induced EPC proliferation and migration. Mechanistically, HGF stimulated Gab1 tyrosine phosphorylation in EPCs, thus leading to activation of extracellular regulated MAP kinase 1/2, which is involved in proliferation and migration signaling.
Up-regulation of Grb2-associated Binder 1 Promotes Hepatocyte Growth Factor Induced Endothelial Progenitor Cell Proliferation and Migration

Qing Fan\textsuperscript{1,2}, Liyu Zhang\textsuperscript{1,2}, Wenjie Zhu\textsuperscript{1,2}, Sheng Xue\textsuperscript{3}, Yisheng Song\textsuperscript{1,2}, Qing Chang\textsuperscript{2}

\textsuperscript{1} Qingdao University, Qingdao 266071, China
\textsuperscript{2} Cardiovascular Surgery Department, The Affiliated Hospital of Qingdao University, Qingdao 266003, China
\textsuperscript{3} Institute for Translational Medicine, Qingdao University, Qingdao 266021, China

Corresponding Author:
Qing Chang
Cardiovascular Surgery Department, The Affiliated Hospital of Qingdao University, Qingdao 266003, China
Email address: changqing20671@163.com

Abstract

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Introduction
Heart and cardiovascular-associated conditions are major causes of death worldwide, and the number of people affected is continually increasing. According to World Health Organization statistics, 17.9 million deaths were due to cardiovascular disease in 2016, accounting for 44% of mortality worldwide (WHO, 2018). Heart valve disease makes up a large proportion of cardiovascular conditions and affects more than 100 million people worldwide (Members et al., 2015).

Currently, mechanical or biological prostheses are the “gold standard” treatment for heart valve failure (Baumgartner et al., 2017). However, both types of valves cannot grow and regenerate — functions particularly important in congenital heart defects, and patients often require subsequent reoperation as they age (Blum, Drews, & Breuer, 2018). All available prostheses have other disadvantages, such as a risk of infection, constant anticoagulation in mechanical valves, progressive degeneration in bioprostheses, and lack of a tissue source in homografts (Best et al., 2016). The tissue-engineering heart valve (TEHV) can potentially overcome most of the current shortcomings by providing a viable valve capable of growth, remodeling, and repair. The ultimate goal of TEHV is to produce a structure similar to the native valve structure (Emmert et al., 2014). In 2006, Cebotari et al. reported two successful clinical applications of TEHV in humans by using autologous endothelial progenitor cells (EPCs) (Cebotari et al., 2006). As one of the main cell sources of TEHV, EPCs can attach, multiply, and cover the implant with an endothelial layer. However, autologous EPCs are usually insufficient in patients who require heart valve replacement, and the covering with an endothelial layer occurs slowly (Bianconi et al., 2018; Rippel, Ghanbari, & Seifalian, 2012). The absence of this autologous endothelial layer may cause immunological reactions. If autologous EPC proliferation and migration could be enhanced, the re-endothelialization time — i.e., the time required for TEHV to form an endothelial layer — could be decreased, and the immunological reactions could be alleviated.

The Grb2-associated binder (Gab) family docking proteins are involved in the amplification and integration of signal transduction evoked by growth factors, cytokines, antigens, and numerous other molecules (Nakaoka & Komuro, 2013). Gab1 has an essential role in postnatal angiogenesis and arteriogenesis via hepatocyte growth factor (HGF)/c-Met signaling. It is associated with SHP2 after stimulation with HGF, which is required for activation of ERK1/2 (Aasrum, Odegard, Sandnes, & Christoffersen, 2013). Downregulation of Gab1 inhibits cell proliferation and migration (Sang, Li, Li, & Liu, 2013; Xu, Li, Kuang, Kuang, & Wu, 2017). In addition, Gab1 is required for HGF-induced EC (endothelial cell) proliferation and migration (Zhao et al., 2011). We hypothesized that Gab1 might play the same role in EPCs, which are EC precursors, and that upregulation of Gab1 in EPCs might have a positive effect on proliferation and migration.

Here, we demonstrated that upregulation of Gab1 promotes HGF-induced EPC proliferation and migration.

Materials & Methods
Isolation and culture of EPCs

Compared with adult peripheral blood or bone marrow progenitors, cord blood progenitors have distinct proliferative advantages, and cord blood can be obtained noninvasively (Murohara et al., 2000). Human umbilical cord blood was collected from the placental cords of volunteers undergoing Cesarean section delivery. Collection occurred immediately after the delivery of the placenta to avoid clot formation. The EPCs were isolated and cultured as described previously (Li et al., 2016; Murohara et al., 2000). Briefly, mononuclear cells were isolated from the human cord blood by density gradient centrifugation over Histopaque-10771 (Sigma-Aldrich, St. Louis, MO, USA), according to the manufacturer’s protocol. The cells were seeded in T25 flasks pre-coated with 0.1 mg/ml of human fibronectin (Sigma-Aldrich) and were incubated in EGM-2 BulletKit medium (Lonza, Cologne, Germany). After 3 days, nonadherent cells were removed, and the medium was replaced. Subsequently, the medium was changed every 2 days.

Characterization of EPCs

After 7 days of culture in vitro, the EPCs were characterized as adherent cells, which were double-positive for acetylated low-density lipoprotein (acLDL) uptake and lectin binding, as assessed by direct fluorescent staining, as described previously.

Briefly, to evaluate the ability of ac-LDL uptake and lectin binding in EPCs, the cells were cultured in 10 μg/mL of 1, 1'-dioctadecyl-3, 3', 3'-tetramethyl-indocarbocyanine perchlorate-labeled acetylated low-density lipoprotein (Sigma-Aldrich) for 4 hours at 37°C. They were then fixed with 2% paraformaldehyde for 15 minutes. The cells were washed with PBS and reacted with fluorescein isothiocyanate (FITC)-labeled Ulex europaeus agglutinin-1 (UEA-1, 10 μg/mL; Sigma-Aldrich) at room temperature for 1 hour. The cells were washed to remove the free UEA-1. Nuclear counterstaining was performed with DAPI (4', 6-diamidino 2-phenylindole; Sigma-Aldrich), and the cells were examined under a fluorescence microscope (Nikon, Tokyo, Japan). The adherent cells that stained with triple-positive fluorescence were determined to be EPCs. Nuclear staining with DAPI verified that nearly all the adherent cells (> 95%) were acLDL (+) ulex-lectin (+).

Adenovirus transfection

AD-Gab1 was purchased from Biowit Technologies (Shenzhen, China). EPCs were harvested with 0.25% trypsin-EDTA (Gibco, Carlsbad, CA, USA) plated and seeded on 96-well plates at a density of 8000 cells per well. After 24 hours, 100 μL AD-Gab1 medium at six different multiplicity of infection (MOI) levels was added into each well for EPC transfection in triplicate. After 24 hours, the cells were examined under a fluorescence microscope, and fluorescence intensities were measured in ImageJ. A suitable MOI was determined and used for subsequent EPC transfection with AD-Gab1. After the cell intensity reached approximately 80%, RNA and protein were extracted from cells. We used a PrimeScript RT Reagent Kit (Takara, Dalian, China) to synthesize cDNA, then examined the expression of Gab1 by using quantitative real-time PCR with SYBR Premix Ex Taq II (Takara), in triplicates run three times each. For Gab1, we used the forward primer 5'-TGCCATTAACTGTGCTTCCCA-3’ and the reverse primer 5’-TCGCACAGAGCACTCCAAAT-3’. For β-actin, we used the forward primer 5’-
CTCCATCCTGGCCTCGCTGT-3’ and the reverse primer 5’-GCTGTCACCTTCACCGTTCC-3’. The relative Gab1 expression was calculated in Bio-Rad CFX Manager. Protein was prepared for western blot analysis in triplicate.

**Measurement of cell proliferation**

EPCs that were cultured at 37°C and 5% CO₂ in an incubator for 14 days were harvested with 0.25% trypsin-EDTA and plated onto 96-well plates with EGM-2. The cells were incubated at 37°C and 5% CO₂ for 24 hours. We added 0.5 μL of 50 μM EdU (Sigma-Aldrich) into each well containing 500 μL of medium and incubated the cells for 4 hours. The cells were fixed with 4% paraformaldehyde and incubated with 2 mg/mL aminoacetic acid for 5 min with oscillation. The cells were incubated with 100 μL of the penetrant into each well and oscillated 10 min, and 100 μL of 1× EdU solution was then added and incubated for 30 minutes. DAPI was used to stain cell nuclei.

After digestion with pancreatic enzyme, EPCs were plated on 96-well plates with a density of approximately 4000 cells per well, in triplicate. The next day (24 hours later), the AD-Gab1 group and AD-Gab1+HGF group were transfected with AD-Gab1 at MOI = 20. One plate was randomly chosen to run CCK-8 (7Sea Pharmatech, Shanghai, China) tests, to determine 24 hour proliferation. The medium of the Control+HGF group and AD-Gab1+HGF group was changed to EGM-2 with 20 ng/mL of HGF (PeproTech, Rocky Hill, NJ, USA), whereas the Control group and AD-Gab1 group were maintained in EGM-2 without HGF. The cells were incubated for another 24 hours, and cell proliferation was then tested (72 hour time point). Subsequently, we performed CCK-8 tests at 96 hours and 120 hours with the last two plates.

**In vitro wound-healing assay**

The cells were removed by trypsinization, counted, and plated at a density of 1×10⁶ cells/well in six-well plates. After transfection, cells were incubated until confluent monolayers were formed for wounding assays. Wounds were made with a pipette tip, and photographs were taken immediately (time 0). Then the medium of the Control+HGF group and AD-Gab1+HGF group was changed to EGM-2 without FBS and with 20 ng/mL of HGF; the Control group and AD-Gab1 group were maintained in EGM-2 without FBS. Photographs were taken at 12 hours, 24 hours, 36 hours, and 48 hours after wounding. The cell-covered area (%) was measured to determine the amount of migration by the cell monolayer to cover the wounded area during this time period. The cell-covered area (%) was determined as (W₀ – Wₜ) / W₀, where W₁ represents the wound area (with no cells) at time t. These areas were measured and analyzed in ImageJ.

**Western blot analysis**

Cells were harvested in lysis buffer. The protein concentration of the lysate was determined by using the Bradford method. Equal amounts of protein were incubated with specific antibody overnight at 4°C with gentle rotation. Then protein A/G PLUS-agarose was added and incubated for an additional 2 hours. Then the beads were washed extensively with lysis buffer, and immune complexes were eluted in SDS-PAGE sample buffer. Total immune complex samples or protein samples from total cell lysate were separated by SDS-PAGE, transferred to a nitrocellulose membrane, and incubated with the appropriate primary antibodies (Cell Signaling Technology,
Danvers, MA, USA). After washing and incubation with secondary antibodies (Cell Signaling Technology), immunoreactive proteins were visualized by using the ECL detection system.

**Statistical analysis**

The original data were processed in GraphPad Prism 7.0. The quantitative real-time PCR data were analyzed with an independent-samples t-test. The western blot analysis in figure 1 was performed with a Student’s t-test. Other statistical analysis was performed with ordinary one-way ANOVA. All these tests were performed on at least three replicates. All data are expressed as the mean ± SD. A value of \( P < 0.05 \) was considered statistically significant.

**Ethics approval and consent to participate**

We have received informed consent from participants. All the procedures were followed by the Medical Ethics Committee of The Affiliated Hospital of Qingdao University, Qingdao.

**Results**

**Characterization of EPCs**

Freshly isolated mononuclear cells were the small round cells suspended in the medium. With increased culturing time, the cells gradually stretched and became larger, forming cell–cell adhesions. Cell colonies were observed at approximately 7 days after seeding. Three to four weeks later, the endothelial cells showed a typical cobblestone-like appearance. After 7 days’ culture in vitro, when observed under a fluorescence microscope, the adherent dil-acLDL-labeled cells were red, the cells bound with FITC-UEA-I were green, and the cell nuclei were stained blue with DAPI. Almost 100% of the cells showed three colors, and these cells were considered the differentiating EPCs (Fig. 1A, B, C, D, E, and F).

**Adenovirus transfection into EPCs**

The efficiency of adenovirus transfection was determined by fluorescence microscopy. The more EPCs transfected, the more green fluorescence was observed. An excessive MOI may damage EPCs and even lead to cell death. The statistical results indicated that, at MOI = 20, the number of cells with green fluorescence was much greater than at other MOI levels (Fig. 1G). After we selected the proper MOI, we sought to validate the adenovirus transfection into EPCs. The qPCR results showed that the expression of the **Gab1** gene was upregulated in the AD-Gab1 group after transfection (Fig. 1H), and western blot analysis demonstrated that Gab1 protein expression was higher in the AD-Gab1 group (Fig. 1I, J).

**Effect of overexpression of Gab1 on EPC proliferation**

Detection of EdU is more sensitive than BrdU detection and can be accomplished in minutes (Salic & Mitchison, 2008). Under a fluorescence microscope, we observed EdU incorporation to determine the proliferation of EPCs that were cultured for 14 days (Fig. 2A, B, C, and D). The proliferation of EPCs in the Control+HGF group and AD-Gab1 group was increased, whereas the proliferation in the AD-Gab1+HGF group was even higher, and the results were statistically significant (Fig. 2E).
In CCK-8 tests, the optical density (OD) was measured at 450 nm wavelength. The greater the cell number, the higher the OD. Because we added HGF stimulation between 48 hours and 72 hours, the cell proliferation rate was determined as the OD value at 72 hours divided by the OD value at 48 hours. Statistical analysis of the data confirmed the results of the EdU incorporation assay. The cell proliferation ability of the Control+HGF group and AD-Gab1 group was higher than that of the Control group between 48 hours and 72 hours (Fig. 2F). The cell proliferation ability of the AD-Gab1+HGF group was significantly different from that of the other three groups (Fig. 2F). We plotted growth curves with the mean OD values as the ordinate and the cell culture time as the abscissa (Fig. 2G). After HGF stimulation, EPCs in the AD-Gab1+HGF group grew most rapidly.

In conclusion, HGF-stimulation and Gab1 have a role in promoting cell proliferation, and upregulation of Gab1 expression enhances this effect.

**Effect of overexpression Gab1 on EPC migration**

In wound-healing assays, we determined the area covered by a wounded cell monolayer on plastic after different treatments. When we used EGM-2 without FBS, EPCs had little proliferation ability. The results are reported as cell-covered area (fig. 3A). The cell-covered areas in the Control+HGF group and AD-Gab1 group were higher than that in the Control group, and the increase in cell-covered area in the Control+HGF group was higher than that in the AD-Gab1 group. The data suggested that HGF stimulation and upregulation of Gab1 both influence EPC migration. HGF stimulation had a greater effect on EPC migration than up-regulation of Gab1 expression. We presumed that HGF stimulation combined with upregulation of Gab1 to treat EPCs should result in a greater improvement in EPC migration. To test this possibility, we analyzed an AD-Gab1+HGF group. At 12 hours, the cell-covered area in the AD-Gab1+HGF group was significantly higher than those in the other three groups (Fig. 3A). At 24 hours, 36 hours, and 48 hours, the cell-covered area in the AD-Gab1+HGF group was higher than those in the other groups (Fig. 3A) and increased more quickly than those in the other groups (Fig. 3B). Thus, EPCs treated with both HGF stimulation and transfection of adenovirus containing the Gab1 gene had a greater migration ability.

**HGF-mediated signaling in EPCs**

The ERK signaling pathway is involved in cell proliferation and migration. To investigate how upregulation of Gab1 mediates HGF-induced EPC proliferation and migration, we assessed the activation of ERK1/2 in response to HGF stimulation and upregulation of Gab1 expression. The expressions of Gab1, phospho-Gab1, SHP2, phospho-SHP2, ERK1/2, and phospho-ERK1/2 in EPCs was analyzed via western blotting. Quantitative analysis demonstrated that the phosphorylation of Gab1 in the AD-Gab1 group was almost equal to that in the Control group but lower than that in the Control+HGF group and AD-Gab1+HGF group (Fig. 4A). We confirmed that HGF stimulated Gab1 tyrosine phosphorylation, as previously reported (Zhao et al., 2011). Once sufficient Gab1 expression was achieved, for example in the AD-Gab1+HGF group, HGF induced more phosphorylation of Gab1 (Fig. 4A). The increased Gab1 phosphorylation substantially increased the levels of SHP2 phosphorylation, thus leading to
activation of MAP kinase 1/2, which is involved in proliferation and migration signaling (Fig. 4B, C). Together, these results established a critical role of EPCs Gab1 in proliferation and migration signaling and indicated that upregulation of Gab1 enhances the function of this signaling pathway.

Discussion

Asahara et al. first reported that a subtype of hematopoietic progenitor cells from adults, namely EPCs (Asahara et al., 1997), have EC (endothelial cell) features and can differentiate into mature ECs and might represent a potential strategy for TEHV. Subsequently, experiments confirmed that TEHV using autologous EPCs is very encouraging (Cebotari et al., 2006; Dohmen et al., 2007; Sales et al., 2010). However, EPCs are a fairly rare cell population, and, when administered intravenously, only a very small fraction of infused cells reach the target region and participate in re-endothelialization. How to increase their cell number is an urgent problem.

Mechanistically, we found that Gab1 is critical in mediating HGF proliferation and migration signaling of human EPCs in vitro. To our knowledge, this is the first study revealing a key role of upregulation of Gab1 in promoting proliferation and migration and HGF-mediated signaling in human EPCs.

Previous studies on the role of Gab1 have usually used a Gab1-ecKO mice model. Those studies have found that Gab1-deficient mice have developmental defects in postnatal angiogenesis (Wang, Xu, Yin, & Jin, 2015; Xu et al., 2017). However, the effect of upregulating Gab1 in human EPCs has not been explored. Using human umbilical cord blood, we isolated human EPCs and upregulated their Gab1 expression through adenovirus transfection. We showed that upregulation of Gab1 markedly enhanced the proliferation and migration of human EPCs. HGF have been shown to mobilize and increase EPC number (Rehman et al., 2004). Our results showed that under the same dose of HGF stimulation, the proliferation and migration of human EPCs was strongly enhanced in EPCs in which Gab1 was upregulated. Therefore, upregulation of Gab1 amplifies stimulation by HGF.

The results also supported that Gab1 is important for HGF-induced ERK1/2 phosphorylation. Few studies in other cells, such as HUVEC and MDCK cells, have evaluated the role of Gab1 in HGF-induced ERK1/2 (Aasrum et al., 2015; Maroun, Naujokas, Holgado-Madruga, Wong, & Park, 2000; Shioyama et al., 2011). In these cells, which were transfected to express mutated Gab1 unable to recruit SHP2, the HGF induced sustained activation of the ERK pathway was found to be reduced (Maroun et al., 2000). ERK1/2 activation has been shown to regulate cell migration and survival signaling pathways (Koch, Tugues, Li, Gualandi, & Claesson-welsh, 2011). Gab1 plays an important role in mediating growth factor-induced activation of ERK1/2 through recruiting SHP2 in a tyrosine phosphorylation-dependent manner (Aasrum et al., 2015). However, the role of Gab1 in HGF mediated signaling in human EPCs remains unclear. Our results suggested that upregulation of Gab1 in human EPCs results in recruitment of more SHP2.
We further found that under the same dose of HGF stimulation, greater SHP2 tyrosine phosphatase activation and ERK1/2 phosphorylation were observed in cells overexpressing Gab1. The data suggest signaling mechanisms by which Gab1 mediates growth factor-induced proliferation and migration.

Conclusions

In this study, we found that upregulation of Gab1 promotes HGF-induced EPC proliferation and migration. Mechanistically, HGF stimulates Gab1 tyrosine phosphorylation in EPCs, thus leading to activation of extracellular regulated MAP kinase 1/2, which is involved in proliferation and migration signaling. Our findings may have clinical implications: they suggest that enhancing Gab1 signaling may be a potential strategy to increase EPC cell number and provide a new means of achieving rapid TEHV re-endothelialization.

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Figure 1

Immunofluorescence staining of EPCs and expression of Gab1 at the optimal MOI.

Fig. 1 Immunofluorescence staining of EPCs was performed after 7 days (orig. mag. ×40). (A) Unstained EPCs; (B) dil-acLDL-labeled cells (red); (C) FITC-UEA-I-bound cells (green); (D) DAPI contrast staining of the cell nuclei; (E) dil-acLDL-labeled and FITC-UEA-I-bound double positive cells; (F) dil-acLDL-labeled, FITC-UEA-I-bound, and DAPI stained triple positive cells. Fluorescence indicating Gab1 expression in EPCs transfected with adenovirus (24 hours) with different multiplicity of infection (MOI) levels, and expression of Gab1 at the optimal MOI. (G) The number of fluorescent cells at different MOI levels. (H) qPCR results show the expression of the Gab1. (I, J) The expression of the Gab1 after transfected (western blot). **p < 0.01 (n = 3) versus other MOI groups, *p < 0.05 (n = 3) versus Control, ***p < 0.001 (n = 3) versus Control. Values are the mean ± SD.
Figure 2

EdU incorporation and CCK-8 cell viability tests reflecting the proliferation of EPCs in different groups.

Fig. 2 EdU incorporation reflecting the proliferation of EPCs cultured for 14 days. (A), (B), (C), (D) DAPI (blue) was used to stain nuclei, and Ed U (green) was incorporated into EPCs in each group (orig. mag. ×100). (E) The ratio of EPCs stained by Ed U in each group. *p < 0.05 (n = 5) versus Control+HGF and AD-Gab1 group, **p < 0.01 (n = 5) versus Control, Control+HGF and AD-Gab1 group. CCK-8 cell viability tests reflecting the proliferation of EPCs in different groups. (F) Cell proliferation in each group. (G) Growth curves of EPCs in different time and treatment conditions. *1p < 0.05 (n = 4) versus Control, Control+HGF and AD-Gab1 group. *2p < 0.05 (n = 4) versus Control+HGF and AD-Gab1 group. Values are the mean ± SD.
Figure 3

Wound-healing assays reflecting the migration of EPCs in different groups.

Fig. 3 Wound-healing assays reflecting the migration of EPCs in different groups. (A) Cell-covered area in each group at different times. (B) Cell-covered area increased over time. * p < 0.05 (n = 3) versus Control, Control+HGF and AD-Gab1 group. Values are the mean ± SD.
Figure 4

Effect of overexpression of Gab1 on HGF-induced phosphorylation of SHP2 and ERK1/2.

The phosphorylation of SHP2 and ERK1/2 was determined by SDS-PAGE followed by blotting with a phospho-specific antibody. Equal loading of each lane was confirmed by reprobing blots for total SHP2 or ERK1/2. Representative blots (left) and bar graphs (right) summarizing the effects of the overexpression of the Gab1 (A) on the HGF-induced (20 ng/mL) phosphorylation of SHP2 (B) and ERK1/2 (C). *P < 0.05 (n = 3) vs Control, Control+HGF and AD-Gab1 group. Values are the mean ± SD.