Lehigh Valley Health Network LVHN Scholarly Works

Department of Obstetrics & Gynecology

Examining the Relationship of Inflammatory Mediators Among Fetal Compartments

Daniel G. Kiefer MD Lehigh Valley Health Network, Daniel G.Kiefer@lvhn.org

Sean M. Keeler MD Lehigh Valley Health Network

Jolene C. Muscat MD

Michael Demishev MD

Nazeeh Hanna MD

Follow this and additional works at: https://scholarlyworks.lvhn.org/obstetrics-gynecology Part of the <u>Obstetrics and Gynecology Commons</u>

Published In/Presented At

Kiefer, D., Keeler, S., Muscat, J., Demishev, M., & Hanna, N. (2011, February 7-12). *Examining the relationship of inflammatory mediators among fetal compartments*. Poster presented at: The 31st Annual Meeting of the Society for Materna-Fetal Medicine, San Francisco, CA.

This Poster is brought to you for free and open access by LVHN Scholarly Works. It has been accepted for inclusion in LVHN Scholarly Works by an authorized administrator. For more information, please contact LibraryServices@lvhn.org.

Examining the Relationship of Inflammatory Mediators Among Fetal Compartments

Daniel Kiefer¹, LCDR Sean Keeler², Jolene Muscat³, Michael Demishev³, Nazeeh Hanna⁴ ¹Lehigh Valley Health Network, Obstetrics and Gynecology, Allentown, PA, ²Naval Medical Center Portsmouth, Obstetrics and Gynecology, Portsmouth, VA, ³Stony Brook-Winthrop University Hospitals, Obstetrics and Gynecology, Long Island, NY, ⁴Winthrop University Hospital, Pediatrics, Mineola, NY

OBJECTIVE: Although cytokines have been shown to be important in both term and preterm labor, the source (ie, fetal or placental) of many cytokines is uncertain. Therefore, we evaluate the relationship of inflammatory mediators between the umbilical artery, umbilical vein, placenta, and amniotic fluid.

STUDY DESIGN: Twenty term, non-laboring patients without major maternal or fetal complications undergoing cesarean delivery were asked to provide samples during the immediate pre-operative and intra-operative period. Amniotic fluid was obtained intra-operatively via needle aspiration of the intact amniotic sac after hysterotomy. Fetal plasma from the umbilical artery and vein were sampled from a section of cord after delivery. A portion of the placenta was cultured under standard conditions for 24 hours. The supernatant was then collected. All fluids were analyzed for 27 inflammatory mediators using the Bio-Plex Array. We the compared the inflammatory mediator profile between compartments using Spearman correlation with P < 0.05 required for significance. Mediator levels in the umbilical artery and vein were compared using the t-test.

RESULTS: Table 1 shows the number of cytokines (out of 27) that reached a significant correlation between the various fetal compartment combinations. There was a high degree of correlation between the umbilical artery and vein with 19 of the 27 reaching a significant correlation. When comparing median cytokine levels, only monocyte chemotactic protein-1 (MCP-1) was significantly different in the artery (196 pg/ml) when compared to the vein (123 pg/ml, P=0.01).

CONCLUSION: There is significant correlation of inflammatory mediators between the umbilical artery and vein, but not between the fetal circulation and amniotic fluid. The contribution of the placenta to the in-utero inflammatory milieu remains unclear as few cytokine levels were correlated with other compartments.

Background and Objective:

- Although cytokines and other inflammatory mediators have been shown to be important in both term and preterm labor, the source (ie, fetal or placental) of many cytokines is uncertain.
- We evaluate the relationship of inflammatory mediators between the umbilical artery, umbilical vein, placenta, and amniotic fluid.

Methods:

- Term, non-laboring patients without major ma undergoing cesarean delivery were asked to immediate pre-operative and intra-operative
- Amniotic fluid was obtained intra-operatively intact amniotic sac after hysterotomy.
- Fetal plasma from the umbilical artery and vei section of cord after delivery.
- A portion of the placenta was cultured under hours. The supernatant was then collected.
- All fluids were analyzed for 27 inflammatory r Plex[™] Array.
- We the compared the inflammatory mediator compartments using Spearman correlation with significance.
- Mediator levels in the umbilical artery and vei non-parametric t-test.

Results:

- Table 1 shows the number of cytokines (out) significant correlation between various fetal
- There was a high degree of correlation between vein with 19 of the 27 reaching a significant
- Table 2 contains the median cytokine levels
- When comparing median cytokine levels, on protein-1 (MCP-1) was significantly different when compared to the vein (123 pg/ml, P=0.

Conclusions:

- There is significant correlation of inflammator the umbilical artery and vein, but not betweer amniotic fluid.
- The contribution of the placenta to the in-utero inflammatory milieu remains unclear as few cytokine levels were correlated with other compartments and there was little change in mediator levels between the umbilical artery and vein.
- It is possible that the culture methods altered placental mediator expression. Future studies will attempt to perform direct analysis of placental tissue.

	aidei	7. Intrammatory Mediator Correlation Among Com				
		Compartment	Fluid	Vein	Artery	Cultu
		Amniotic fluid		1	1	2
		Umbilical Vein			19	5
		Umbilical Artery				
		Placental Culture				
		Table indicates the number	r of inflommatory n	podiatora (out of t	07) that reached a	oignificant o
	betw	veen the indicated compartm	nents.		27) that reached a	a significant c
	Table 2.	Median Inflam	matory Me	ediator Le	evels by C	ompart
		Compartment	Amniotic	Umbilical	Umbilical	Placenta
rd conditions for 24				Vein	Vein	
		IL-1ra	930.3	167.6	104	239.7
rs using the Bio-		IL-2	Not detected	Not detected	Not detected	Not deter
		IL-4	1.8	1.1	0.7	4.1
		IL-5	3.4	0.9	0.6	0.9
.05 required for		IL-6	25.3	3.0	3.6	00R>
		IL-7	12.6	2.7	3.7	4.2
		IL-8	40.0	4.7	4.4	29855
compared using the		IL-9	21.3	7.6	4.5	48.9
		IL-10	1.9	1.9	1.7	239.
		IL-12	13.0	6.1	4.6	5.9
		IL-13	5.1	2.4	2.2	53.6
		IL-15	Not detected	8.0	7.5	21.6
		IL-17	9.9	27.5	23.1	56.1
tions.		PDGF	37.1	241.8	257.4	93.8
and		Eotaxin	Not detected	45.6	51.8	20.3
		FGF	13.7	46.0	26.3	181.2
		G-CSF	99.4	43.7	42.9	00R>
			Not detected	100.2	100.2	263.7
			300.4	1976	00.5 272 9	16209
		MCP-1	28.2	190.2	97.7	12803
		MIP-1A	Not detected	Not detected	Not detected	5451
		MIP-1B	6.2	81.4	84 1	13838
		RANTES	22 5	1001 7	943.8	1178
		ΤΝΑ-α	Not detected	18.0	16.1	4044
		VEGE	Q 11	30.3	25 /	
	KEY: Interleuk (G-CSF), granu chemotactic pr	tin (IL), platelet derived grown locyte macrophage stimulati otein-1 (MCP-1), macrophag	th factor (PDGF), fill ng factor (GM-CSF le inflammatory pro	broblast growth fa), interferon gami otein (MIP), tumor	actor (FGF), grant ma (IFN-γ), induci necrosis factor a	ulocyte colony ble protein-10 Ipha (TNF-α),

A PASSION FOR BETTER MEDICINE."

ents

(pg/ml)

ng factor onocyte ndothelia *Out of range, greater than the limit of the assay.

Disclaimer: The views expressed on this poster are those of the author and do not necessarily reflect the official policy or position of the Department of the Navy, Department of Defense, or the U.S. Government.

