AD_____

Award Number: W81XWH-12-1-0611

TITLE: Metabolic Signature of Antipsychotics used in the Treatment of Autism".

PRINCIPAL INVESTIGATOR: Nira Ben-Jonathan

CONTRACTING ORGANIZATION: University of Cincinnati Ôa & a } action of Cincinnati

REPORT DATE: October 2013

TYPE OF REPORT: 05; } * æ

PREPARED FOR: U.S. Army Medical Research and Materiel Command Fort Detrick, Maryland 21702-5012

DISTRIBUTION STATEMENT: Approved for Public Release; Distribution Unlimited

The views, opinions and/or findings contained in this report are those of the author(s) and should not be construed as an official Department of the Army position, policy or decision unless so designated by other documentation.

REPORT DOCUMENTATION PAGE					Form Approved	
			-		OMB No. 0704-0188	
Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing this collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to Department of Defense, Washington Headquarters Services, Directorate for Information Operations and Reports (0704-0188), 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302. Respondents should be aware that notwithstanding any other provision of law, no person shall be subject to any penalty for failing to comply with a collection of information if it does not display a currently valid OMB control number. PLEASE DO NOT RETURN YOUR FORM TO THE ABOVE ADDRESS.						
1. REPORT DATE	2	2. REPORT TYPE			DATES COVERED	
October 2013		Annual			September2012-29September2013	
4. TITLE AND SUBTIT		and in the Treation		5a.	CONTRACT NUMBER	
Metabolic Signatur	e of Antipsychotics	used in the Treatm	ent of Autism .			
					31XWH-12-1-0611	
				5c.	PROGRAM ELEMENT NUMBER	
6. AUTHOR(S) Nira Ben-Jonathan				5d.	PROJECT NUMBER	
				5e.	TASK NUMBER	
				5f.	WORK UNIT NUMBER	
7. PERFORMING ORG	ANIZATION NAME(S)	AND ADDRESS(ES)		8. 1	PERFORMING ORGANIZATION REPORT	
University of Cincir				1	NUMBER	
Cincinnati, OH 452	21-0222					
		AME(S) AND ADDRES	2(52)	10	SPONSOR/MONITOR'S ACRONYM(S)	
	Research and Mat		5(23)	10.	SPONSOR/MONITOR S ACRON HM(S)	
Fort Detrick, Maryl						
T OT Detrick, Maryn				11.	SPONSOR/MONITOR'S REPORT	
					NUMBER(S)	
12. DISTRIBUTION / A	VAILABILITY STATEM	IENT				
Approved for Publi	c Release; Distribu	tion Unlimited				
13. SUPPLEMENTARY NOTES						
14. ABSTRACT						
Background: Atypical antipsychotics (AAP) are prescribed to numerous autistic patients to treat symptoms of agitation,						
stereotypic behavior, temper tantrums and self-injury. Despite their ability to ameliorate many behavioral problems, AAP have						
serious metabolic side-effects which include weight gain, insulin resistance, and increased risk of diabetes and cardiovascular						
disease. The main therapeutic targets of AAP are the dopamine (DAR) and serotonin (5-HTR) receptors. The general						
consensus is that AAP cause metabolic disturbances by an exclusive action on the brain. Preliminary Data: We discovered						
functional DAR and 5-HTR subtypes in human adipose tissue and found that incubation of adipose explants and adipocytes						
with olanzapine, risperidone and ziprasidone suppressed leptin and adiponectin and alter inteleukin-6 (IL-6) release. Oral						
delivery of olanzapine to female rats caused a rapid and robust suppression of leptin, a satiety hormone, concomitant with						
increased food intake and weight gain. Hypothesis and Objectives: We hypothesized that activation of DAR and/or 5-HTR						
subtypes in adipose tissue contributes to the metabolic side-effects caused by AAP. The overall objective was to establish						
adipose tissue as a critical target of AAP and elucidate some of the mechanisms by which the drugs alter adipose tissue						
functions leading to weight gain and the metabolic syndrome.						
15. SUBJECT TERMS- none provided						
16. SECURITY CLASS	IFICATION OF:		17. LIMITATION OF ABSTRACT	18. NUMBER OF PAGES	19a. NAME OF RESPONSIBLE PERSON USAMRMC	
a. REPORT	b. ABSTRACT	c. THIS PAGE	1		19b. TELEPHONE NUMBER (include area	
U	U	U	UU		code)	
			_			

Table of Contents

<u>Page</u>

Introduction	3
Body	3
Key Research Accomplishments	5
Reportable Outcomes	5
Conclusion	5
References	5
Appendices	6

Introduction

Background: Atypical antipsychotics (AAP) are prescribed to numerous autistic patients to treat symptoms of agitation, stereotypic behavior, temper tantrums and self-injury. Despite their ability to ameliorate many behavioral problems, AAP have serious metabolic side-effects which include weight gain, insulin resistance, and increased risk of diabetes and cardiovascular disease. The main therapeutic targets of AAP are the dopamine (DAR) and serotonin (5-HTR) receptors. The general consensus is that AAP cause metabolic disturbances by an exclusive action on the brain.

Preliminary Data: We discovered functional DAR and 5-HTR subtypes in human adipose tissue and found that incubation of adipose explants and adipocytes with olanzapine, risperidone and ziprasidone suppressed leptin and adiponectin and alter inteleukin-6 (IL-6) release. Oral delivery of olanzapine to female rats caused a rapid and robust suppression of leptin, a satiety hormone, concomitant with increased food intake and weight gain.

Hypothesis and Objectives: We hypothesized that activation of DAR and/or 5-HTR subtypes in adipose tissue contributes to the metabolic side-effects caused by AAP. The overall objective was to establish adipose tissue as a critical target of AAP and elucidate some of the mechanisms by which the drugs alter adipose tissue functions leading to weight gain and the metabolic syndrome.

Specific Aims:

Specific Aim 1: To determine whether weight-inducing AAP stimulate adipogenesis, enhance lipid accumulation and/or alter expression and release of selected adipokines in human and rat adipocytes *in vitro*.

Specific Aim 2: To examine whether drug-induced leptin suppression is a major drive for increased appetite and weight gain in a rat model.

<u>Body</u>

1. Responsiveness of rat sc and vis fat depots to different ligands and AAP

Although AAP bind primarily to DAR and 5-HTR (1), they also affect histamine receptors (2). In addition, adipose tissue from visceral (vis) and subcutaneous (SC) depots have different properties and are likely to respond differentially to the drugs given a dissimilar distribution of receptors as well as their coupling to signal transduction pathways (3,4). Thus, the first experiment was designed to compare the responsiveness of vis and sc explants from rats to the different ligands and AAPs alone, or in combination. As key endpoints, we selected two key adipokines: leptin and adiponectin (adipo), and two key transcription factors: PPARG, which regulates adipogenesis, and SREBP1, which regulates lipid homeostasis. Periovarian (vis) and sc fat from untreated female rats were

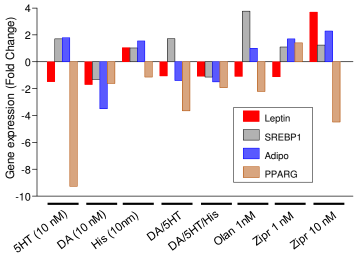


Fig 1: Changes in gene expression in rat sc adipose explants incubated with various compounds.

cut into explants and incubated with serotonin (5HT, 10 nM), dopamine (DA, 10 nM), histamine (His, 10 nM), or a combination of DA/5HT, and DA/5HT/His, each at 10 nM. In addition, explants were incubated with olanzapine (Olan, 1nM and 10 nM) and ziprasidone (Zipr, 10 nM). After 3 days, expression of the four genes was determined by qPCR.

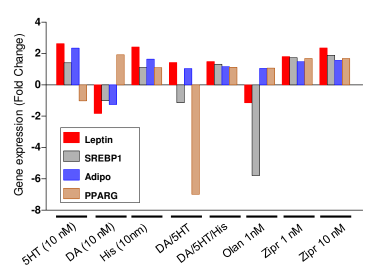


Fig 2: Changes in gene expression in rat vis adipose explants incubated with various compounds.

was slightly stimulated by 5HT, His and Zipr. Although 5HT or DA alone showed no, or slight stimulatory effects PPARG. on their combination showed a marked inhibition. Unlike its stimulatory effect on SREBP1 in sc fat, olanzapine had a significant stimulation in vis fat. Another difference was noted with respect to leptin, whereby 5HT increased leptin expression in vis fat but inhibited its expression in sc fat. The conclusions from these experiments are: 1) there are clearly direct effects of DAR and 5HTR ligands, as well as AAP on both vis and sc fat, 2) histamine receptors do not play a decisive role, 3) a balance between DAR and 5HTR in each depot dictates the overall effects on selected adipose-related genes.

As evident in **Fig 1**, His alone or in combination had little effect on the expression of these genes in sc fat. PPARG was markedly suppressed by 5HT and Zipr, and to a lesser extent by 1 nM Olan or a combination of DA/5HT, while SREBP1 was stimulated by 1 nM Olan. Leptin was increased by Zipr and slightly suppressed by 5HT and DA, while Adipo was slightly stimulated by 5HT and Zipr and DA. significantly suppressed by The combination of DA/5HT did not result in synergism, but in smaller effects.

A rather different profile of gene expression was seen in vis explants (**Fig 2**). PPARG and SREBP1 were markedly suppressed by DA/5HT and 1 nM Olan, respectively, whereas Leptin

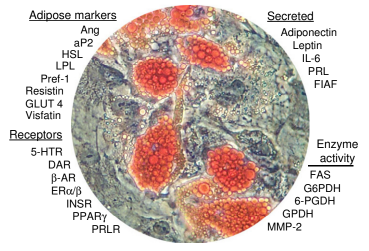


Fig 3: Characterization of LS14 cells. *Left:* expression of adipocyte markers, adipokines and receptors by RT-PCR. *Right:* secreted hormones and enzyme activities.

2. Validation of LS14 cells as representative of vis adipocytes

Although many of our proposed experiments can be done with primary adipocytes, there are

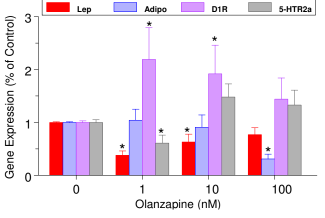


Fig 4: Olanzapine differentially suppresses leptin (Lep), adiponectin (Adipo) and 5-HTR2a, while stimulating D1R expression in LS14 cells, as determined by qPCR.

several caveats, including limited availability and patient-to-patient variability. On the other hand, the advantage of an adipocyte cell line is homogeneity, availability and ease of genetic/biochemical manipulation. Thus, we opted to carry out full characterization of our LS14 cells, which we have cloned from a patient with metastatic liposarcoma (5). Fig 3 shows lipid storage in differentiated LS14 cells stained with Oil-red O. We have used multiple methods such as RT-PCR, Western blotting, ELISAs and enzyme assays, to extensively characterize these cells and establish that they resemble vis adipocytes.

We next examined whether Olanzapine alter the

expression of leptin, adiponectin, D1R and 5-HTR2a in differentiated LS14 cells. Cells were incubated with increasing doses of olanzapine and ziprasidone for 6 hrs, followed by quantitative PCR. (qPCR). Olanzapine at 1 nM inhibited leptin and 5-HTR2a, but stimulated D1R expression, whereas adiponectin was suppressed only at the high dose (**Fig 4**); ziprasidone was less effective than olanzapine (not shown). Unlike 3T3-L1 adipocytes which downregulate leptin expression, LS14 cells produce significant amounts of leptin. The similar responsiveness of LS14 cells and primary adipocytes to the drugs validated the use of either cell type for studying the metabolic effects of AAP *in vitro*.

Key Research Accomplishments

- Establishing the differential responsiveness of vis and sc fat to various agonists and AAP.
- Demonstrating a direct effect of AAP on the expression of critical adipose-related genes.
- Establishing the resemblance of the LS14 adipocyte cell line to visceral adipocytes and validating its use for these studies.

Reportable Outcome

Presentations in Scientific Meetings:

- Ben-Jonathan: Antipsychotic-induced Obesity, BIT's Major Disease Clinical Summit, Warsaw, Poland, November 2013 (Appendix 1)
- Ben-Jonathan: Antipsychotic induced obesity: Direct actions on the adipocytes, EuroSciCon, Anti-obesity drug discovery and development, London, April 2014 (Appendix 2).

Conclusion

We are now well positioned to proceed with a a more comprehensive investigation of the effects of AAP on adipogenesis, enhance lipid accumulation and/or alter expression and release of selected adipokines in human and rat adipocytes *in vitro*.

References

- 1. Nasrallah HA 2008 Atypical antipsychotic-induced metabolic side effects: insights from receptor-binding profiles. Mol Psychiatry 13:27-35
- Han M, Deng C, Burne TH, Newell KA, Huang XF 2008 Short- and long-term effects of antipsychotic drug treatment on weight gain and H1 receptor expression. Psychoneuroendocrinology 33:569-580
- 3. Fain JN, Madan AK, Hiler ML, Cheema P, Bahouth SW 2004 Comparison of the release of adipokines by adipose tissue, adipose tissue matrix, and adipocytes from visceral and subcutaneous abdominal adipose tissues of obese humans. Endocrinology 145:2273-2282
- 4. Borcherding DC, Hugo ER, Idelman G, Richtand NW, Loftus J, Ben-Jonathan N 2011 Dopamine receptors in human adipocytes: expression and functions. PLoS One 6:e25537
- 5. Hugo ER, Brandebourg TD, Comstock CE, Gersin KS, Sussman JJ, Ben-Jonathan N 2006 LS14: a novel human adipocyte cell line that produces prolactin. Endocrinology 147:306-313

BIT Congress-Europe

BIT's Major Diseases Clinical Summit-2013



Dr. Nira Ben-Jonathan, Professor, University of Cincinnati, USA

Speech Title: Antipsychotics-Induced Obesity

Speech Session: 4-2

Highlight of Your Speech: (5-6 Points)

- Atypical antipsychotics induce weight gain and the metabolic syndrome
- Adipocytes serve as a major target for the antipsychotics
- Drug-induced suppression of leptin results in increased food intake
- Suppression of adiponectin contributes to the metabolic syndrome
- Human adipocytes can be used to screen for new drugs devoid of metabolic side effects

Abstract:(within 200 words)

Atypical antipsychotics (AAP) are prescribed to millions of patients with schizophrenia, bipolar disorder, major depression, and autism. Although AAP ameliorate mental dysfunctions, they have serious metabolic side-effects such as weight gain, diabetes, and cardiovascular disease. The primary therapeutic target of the antipsychotics are dopamine (DAR) and serotonin (5-HTR) receptors. The mechanisms underlying the metabolic side effects of AAP have been attributed to their central action. We discovered expression of functional DAR and 5-HTR subtypes in human adipocytes. Incubation of adipose explants and adipocytes with selected AAP suppressed leptin and adiponectin, and increased lipolysis. Treatment of rats with olanzapine caused marked suppression of leptin and adiponectin, and an increase in interleukin-6 (IL-6) expression in fat tissue within 24 hrs, concomitant with increased food intake and weight gain in 2-3 days. We propose that direct activation of DAR and possibly 5-HTR subtypes in adipose tissue by AAP contributes to weight gain and the metabolic syndrome. Human adipocytes could be integrated into the screening paradigm of candidate new drugs for the identification of undesirable metabolic activities prior to costly animal studies and clinical trials. The long term goal is to provide safer drugs to patients requiring treatment with such medications.

Biography:(within 150 words)

Nira Ben-Jonathan, Ph.D, is Professor of Cancer and Cell Biology at the University of Cincinnati, Ohio, USA. She published over 160 manuscripts, edited one book, and contributed 12 chapters to textbooks and encyclopedias. Early in her career she conducted research on the neuroendocrine regulation of pituitary functions. More recently, the focus of her research has shifted to breast cancer and human obesity. Throughout her career, she mentored 65 students, postdoctoral fellows, research scientists and assistant professors. She served on may journal editorial boards and committees of scientific societies. She was awarded the NIH Research Career Development Award, was elected Fellow of the AAAS, was elected Chairman of the Gordon Research Conference, and received the prestigious Rieveschl Award for Outstanding Scientific Research. She has been a member and chairman on numerous study sections of the NIH, the Komen foundation and the DOD.



EuroSciCon Speaker's Form

Please fill in within a month of receipt

*A preliminary title will do, this can be changed up to 2 weeks before the meeting - Thank you

Title of the meeting you are speaking at	Anti-obesity drug discovery and development		
Title (Mrs, Ms, Miss, Mr, Dr, Other)dx	Dr.		
First Name and Surname	Nira Ben-Jonathan		
Job Title	Professor		
Affiliation (Company/Institution/Hospital/University)	University of Cincinnati		
Do you have a website that we can refer to on our site? What is the url?	No		
* Talk Title	Antipsychotics induced obesity: Direct actions on the adipocytes		
* Brief Abstract of your Talk (100 words maximum). This will be place on the meeting agenda.	Atypical antipsychotics (AAP) are prescribed to millions of patients with mental diseases. Although AAP ameliorate mental dysfunctions, they have serious metabolic side-effects such as weight gain, diabetes, and cardiovascular disease. We discovered expression of functional dopamine and serotonin receptors in human adipocytes and found that AAP altered many of their functions. We propose that direct actions of AAP on adipose tissue contribute to weight gain and the metabolic syndrome. Human adipocytes could be integrated into the screening paradigm of candidate new drugs for the identification of undesirable metabolic activities prior to costly animal studies and clinical trials.		
Up to 5 keywords relevant to your talk	adipocytes, metabolic syndrome, antipsychotics, gene expression, adipokines		
Short Professional Biography (100 words maximum) This will be used for promotional purposes and may be used in any press releases about this event .It will also be placed on the meetings web site, the agenda and may be published in our meeting report. Please could you write this in the 3 rd person	Nira Ben-Jonathan, Ph.D, is Professor of Cancer and Cell Biology, University of Cincinnati, Ohio, USA. She published 160 manuscripts, edited one book, and contributed 12 chapters to textbooks and encyclopedias. The focus of her research is on the regulation of pituitary functions, breast cancer and human obesity. She mentored 65 students and scientists, served on journal editorial boards and scientific committees, and has been a member and chairman on NIH, DOD and Komen study sections.		



She was elected Fellow of the AAAS and Chairman of the Gordon Research Conference, and received the Rieveschl Award for Outstanding Research.

*A preliminary title and description will do, this can be changed up to 2 weeks before the meeting - Thank you

Speaker Copyright:

All speaker presentations placed on EuroSciCon computers on the day of the event remains the property of the author, and will be removed from Euroscicons equipment within 24 hours of the event

General Indemnity

EuroSciCon reserves the right to change meeting content, timing, speakers or venue without notice. The event may be postponed or cancelled due to acts of terrorism, war, extreme weather, industrial action, acts of God or any event beyond the control of EuroSciCon. If a situation arises we may endeavour to reschedule the event. EuroSciCon cannot be held responsible for any cost, damage or expenses which may be incurred by the customer as a consequence of the event being postponed. Expenses for speakers are paid only if they are agreed and authorised by EuroSciCon before confirmation of participation by the speaker and meet the criteria as stated on the expense claim form.

Any other comments

Please email to melissa.fletcher@euroscicon.com