Antibiotics Resistance and Integron Class 1 among Commonsal *Escherichia coli*

Manal M. khadhim ¹ and Meraim A. Kazaal ²

Address: ¹ Department of Medical Microbiology, College of Medicine, Al-Qadisiya University, Diwaniya, Iraq, Email: mic_drmanal@yahoo.in .² Department of medical microbiology, College of Medicine, Al-Qadisiya University, Diwaniya, Iraq, Email: medmeraim88@gmail.com

الخلاصة

الأحياء المجهرية المعوية المتعايشة هي مخازن للجينات المقاومة للمضادات الحياتية خصوصا البكتريا القولونية (أي كولأي). وكان الهدف من هذه الدراسة هو التعرف على مدى انتشار الانتكرون الصنف 1 و المقاومة للمضادات الحياتية بين عز لات الأي كولاي المتعايشة. وكذلك، هذه الدراسة تهدف للكشف عن علاقة الانتكرون الصنف 1 مع المقاومة للمضادات الحياتية (مقاومة لمضاد حياتي واحد أو اكثر) و التي قد تنقل أفقيا الى بكتريا أخرى . ثلاثة مائة و واحد عُزلة من الاي كولاي تم الحصول عليها من أشخاص أصحاء (أعمار هم من 1-80 سنة). تم اختبار حساسية جميع ألعز لات ضد 16مضاد حياتي باستخدام طريقة الانتشار القرصي وتم الكشف عن الانتكرون بواسطة PCR. اكتشف الانتكرون الصنف 1 في 112 عزلة (37٪) من أصل 301 عزلة. ولاحظ مقاومة عالية لمعظم المضادات الحياتية خصوصا الامبسلين و الاموكسلين و لينكومايسين و سيفالكسين في حين كانت المقاومة لأميكاسين والجنتاميسين أقل شيو عا. المقاومة لسيفوتاكسيم و سيبروفلوكساسين و أوكسى تتراسكلين والتتراسيكلين وميثوبريم ترتبط بشكل واضح (P<0.05) مع العزلات الحاوية على الانتكرون الصنف 1. كل العزلات الحاوية على الانتكرون الصنف 1 كانت مقاومة لأكثر من خمسة مضادات حياتية ومرتبط بشكل كبير مع العديد من أنماط المقاومة. في الختام، از دياد مقاومة المضادات الحياتية بين عز لات الأي كولاي المتعايشة في هذا المجتمع وكذلك ازدياد انتشار الانتكرون الصنف 1والذي مرتبط بو ضوح مع المقاومة لمضاد حباتي و أحد أو أكثر

Abstract

Normal intestinal microbiota are a reservoir for antibiotic resistance (genes) especially *Escherichia coli* (*E. coli*). The goal of this study was to investigate the prevalence of integrons class 1 and antibiotic resistance among commensal *E. coli* isolates and detect the association of integron class 1 with antibiotic resistance (single or multiple drug resistance) that may be transport horizontally to other bacteria. Three hundred one *E. coli* isolates from stools were obtained from healthy individuals (age from 1to 80 years). All isolates were tested for their susceptibility against 16 antimicrobial agents using standard disc diffusion method and for the presence of class 1 integrons by PCR. Integrase genes detected in 112 isolates (37%) out of 301 tested isolate. High resistance observed for

most antibiotics especially ampicilln, amoxcillin, lincomycin, cephalexin while resistance to amikacin and gentamicin was less common. Furthermore resistance to cefotaxime, ciprofloxacin, oxytetracyclin, tetracycline and trimethoprim significantly associated (p < 0.05) with *Intl1* gene positive isolates. All integrons positive isolates are resist to more than five antibiotics and significantly associated with many resistance patterns. In conclusion, increase antibiotic resistance among commensal *E. coli* in this population that companied with increase frequency of integron class 1 that significantly associated with resistance for single or multiple antibiotic agents.

Key words: Commensal *E. coli*, integron class 1, antibiotic resistance, multiple drug resistance

Introduction

The emergence of antibiotic resistance among pathogenic and commonsal bacteria became one of most clear global health challenges of this century (1). Resistance to antibiotics has a high predominant in bacterial isolates from developing countries which are result from over and improper use of antimicrobial agents. Commensal E. coli inhabit the intestine of many mammals including human and act as a potential reservoir for antimicrobial resistance genes and play an important role in the ecology of antimicrobial resistance of bacterial populations. The prevalence of resistance in commensal E. coli is a useful indicator of antibiotic resistance in bacteria in the community (2,3). Commensal E. coli strains efficiently exchange genetic material with other pathogens such as Salmonella, Shigella, Yersinia and Vibrio, as well as pathogenic E. coli (2,4,5). Recently, exchange of many different and diverse genes responsible for antibiotic resistance has been linked to genetic structures called integrons, that integrate and mobilize individual gene cassettes encoding antibiotic resistance determinants (6). Integron that have resistance gene can be moved to other genetic sites or transferred horizontally to other bacteria mainly by conjugative plasmid. There are many types of integron have been identified and distinguished by their respective integrase (*IntI*) genes (1). Class 1 integrons are strongly

associated with multi-resistance seen in enterobacteriaceae in the hospital environment (7). The present study aimed to indicate the role of commensal *E. coli* as reservoir of single or multiple antibiotic resistance that can be transported by potential transposable elements as integron to pathogenic and commonsal strains in human gut.

Materials and Methods

Study design and sample collection: a cross sectional study that come in agreement with ethics of Al-Diwaniya Teaching Hospital and verbal informed consent was obtained from all healthy participants. Fecal samples collected from healthy workers, officers, students and their families seen in College of Medicine of Al-Qadisiya University and from clinical staff and their family of Al-Diwaniya Teaching Hospital from march 2016 to may 2016.

Isolation and Identification of Microorganisms: The identification of *E. coli* was done depending on morphological features and the rose pink color of the colonies on MacConkey agar plats that confirmed by subculture on Eosin Methylene Blue agar and incubated for 24 hours at 37°C, the typical greenish metallic sheen color indicate of *E. coli* (24) then the result confirmed by biochemical tests and microscopically examination with Gram's stain. After primary identification of *E. coli* bacterial cell cultured in Nutrient broth for DNA extraction for molecular study and cultured on Muller- Hinton agar for antibiotic susceptibility test.

Antibiotic Susceptibility Test: All identified *E. coli* isolates were subjected to sensitivity test by disc-diffusion method (25). Antibiotic resistance of 16 antibiotic discs (ampicillin, amoxicillin, amikacin, norfloxacin, streptomycin, trimethoprim, lincomycin, gentamicin, ciprofloxacin, nalidixic acid, chloramphenicol, cefotaxime, ceftriaxone, cephalexin, tetracyclin and oxytetracyclin) was determined according to the guidelines recommended by the CLSI (2013), corresponding to the drugs considered routine testing and reporting on *Enterobacteriaceae* (25).

Molecular study: Conducted by conventional PCR for amplification primer of conserved region of *IntI1* gene (forward: GGTCAAGG ATCTGGATTTCG and reverse: ACATGCGTGTAAATCATCGTC) (26). Statistical analysis: Statistical analyses were performed using the odd ratio and χ^2 test or Fisher's exact test. A P -value of <0.05 was considered to be statistically significant. Statistical analyses were performed by the Statistical Package for Social Sciences version 17 for Windows software and Microsoft Excel 2010.

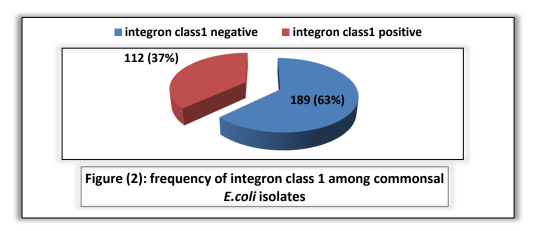
Results

Frequency Of Integron Class 1 and Antibiotic Resistance Among Commonsal E. coli Isolates

Integrase genes detected in 112 isolates (37%) out of 301 tested isolate by PCR amplification (figure 1 and 2).



Figure (1): PCR amplification of IntI1gene of E. coli flora



Among the drugs under the study, ampicillin, amoxicillin, cephalexin and lincomycin not have any antimicrobial effect (100% of the total isolates were resistant). In addition, the highest resistance rate was recorded to cefotaxime (93%), tetracyclin (74%), trimethoprim and oxytetracyclin (67%), ceftriaxone (63%), nalidixic acid (56%) and high resistance also investigated for streptomycin (53%). on the other hand low resistance indicated to gentamicin (21%), ciprofloxacin (26%), norfloxacin (31%) and chloramphenicol (35%) while resistance to amikacin was less common and seen in only 14% of the isolates (table 1). Compared resistance antibiotic of integrons positive $E.\ coli$ isolates with integronsnegative $E.\ coli$ isolates in table 1 reflect that resistance to cefotaxime, ciprofloxacin, oxytetracyclin, tetracycline and trimethoprim are significantly associated with IntII gene positive isolates (P=0.0001).

Table (1): association between antibiotic resistance and integron class 1

Antibiotic agents	Total resistance N (%)	IntI1 positive isolates (N=112) R N (%)	\mathbf{X}^2	OR	<i>P</i> value					
Pencillines										
Ampicillin	301(100)	112(100)	0.075	1.44	0.61					
Amoxcillin	301(100)	112(100)	0.075	1.44	0.61					
Cephalosporin										
Cefotaxime	280(93)	112(100)	33.14	4.44	0.0001					
Ceftriaxone	188(63)	77(69)	2.60	1.02	0.111					
Cephalexin	301(100)	112(100)	0.075	1.44	0.61					
Quinolones										
Nalidixic acid	168(56)	69(62)	1.77	1.36	0.076					
Norfloxacin	92(31)	35(31)	0.033	1.17	0.822					
Ciprofloxacin	77(26)	21(19)	17.40	4.45	0.0001					
Aminoglycsides										
Amikacin	42(14)	17(15)	0.302	1.39	0.333					
Gentamicin	64(21)	26(23)	0.109	1.25	0.544					
Streptomycin	161(53)	63(56)	0.331	1.26	0.491					
Tetracyclines										
Oxytetracyclin	201(67)	95(85)	60.32	7.99	0.0001					
Tetracyclin	223(74)	99(88)	43.66	6.87	0.0001					
Phenicol										
Chloramphenicol	105(35)	44(39)	2.998	1.93	0.1310					
Anti -Folate										
Trimethoprim	202(67)	86(77)	17.77	2.35	0.0001					
lincosamid	_									
Lincomycin	301(100)	112(100)	0.075	1.44	0.61					
Pearson Correlation Coefficient (r) 0.9766										

N = number, R= resistance

Association Between Multiple Antibiotic Resistance And Integron Class 1 among Commonsal Escherichia coli

In present study, all integrons positive isolates are resist to more than five antibiotics (table 2). Among *IntI1* gene positive isolates resist to 6 (9%), 8 (7%), 9 (8%), 10 (16%), 11 (24%), 12 (12%), 13 (12%), 14(9%) and 15 (4%) antibiotics are observed but significantly association (p < 0.05) demonstrated to resistance of 10, 11 and 12 antibiotics only.

Table (2): Comparsion multiple antibiotic resistance between integron positive and integron negative isolates

Number of antibiotics	IntI1 Positive isolates (N=112) R N(%)	IntI1 negative isolates (N=189) R N(%)	P value
4	0(0)	17(9)	NS
5	0(0)	11(6)	NS
6	10(9)	44(23)	NS
8	8(7)	23(12)	NS
9	9(8)	34(18)	NS
10	18(16)	0(0)	< 0.05
11	27(24)	11(6)	< 0.05
12	13(12)	11(6)	< 0.05
13	13(12)	23(12)	NS
14	10(9)	11(6)	NS
15	4(4)	4(2)	NS

R= Resist; N = Number; NS= Non-Significant (P > 0.05)

Many resistance patterns are seen among integrons positive isolates and most common resistance patterns are AM-AX-L-CL (100%) and AM-AX-L-CL-CTX (100). Integron class1 positive isolates are significantly associated (P < 0.05) with all detected resistance patterns in table 3 except AM-AX-L-CL (P = 0.655), AX-AM-CTX-CL-TE-T-CIP-NOR-AN-TMP-C-L-AK-CN-CRO (P = 0.103), AX-CTX-T-C-CRO-S-TE-

CL-TMP-NA-L (P = 0.076), NA-CTX-CIP-T-C-CRO-S-TE-NOR-TMP (P = 0.077) and AX-CTX-T-C-CRO-S-TE-CL-TMP-AM-L (P= 0.076).

Table (3): Distribution of MDR among IntI1 positive isolates

Resistance patterns	Integron class 1-positive isolates (N=112)		Integron class 1- negative isolates (N=189)		OR	X ²	P value	
	R		R					
	N	%	N	%				
AM-AX-L-CL	112	100	189	100	0.0812	1.99	0.655	
AM-AX-L-CL-CTX	112	100	157	83	42.014	34.99	< 0.0001	
AM-AX-L-CL-CTX-TE	99	88	100	53	6.22	43.811	< 0.0001	
AM-AX-L-CL-CTX-TE-NA	62	55	55	29	3.112	19.61	< 0.0001	
AM-AX-TE-T	95	85	55	29	30.61	90.16	< 0.0001	
NOR-CIP-NA	17	15	45	24	0.931	4.95	0.041	
C-TMP-L	39	35	34	18	2.653	9.191	0.006	
CTX-CRO-CL	73	65	89	47	3.304	12.67	0.0021	
NOR-CIP-NA-CTX-CRO-CL	17	15	23	12	12.33	15.73	0.00032	
AX-NA-CTX-T-CL-TMP-AM-L	56	50	55	29	1.19	14.20	0.0001	
AX-CTX-T-TE-S-CL-TMP-AM-L	39	35	45	24	1.39	4.009	0.043	
CRO-TE-CL-TMP	60	54	55	29	2.997	19.91	< 0.0001	
AX-AM-CTX-CL-TE-T-CIP-NOR-AN- TMP-C-L-AK-CN-CRO	5	4	11	6	0.711	0.686	0.103	
AX-CTX-T-C-CRO-S-TE-CL-TMP- NA-L	13	12	11	6	6.05	3.44	0.076	
AX-CTX-T-CRO-TE-CL-TMP-NA-L	43	38	34	18	2.11	11.60	0.0004	
AX-CTX-T-CRO-TE-CL-TMP-L	60	54	34	18	5.26	39.17	< 0.0001	
NA-CTX-CIP-T-C-CRO-S-TE-NOR- TMP	5	4	11	6	0.607	3.44	0.077	
NA-CTX-T-CRO-S-TE-TMP	26	23	34	18	1.35	1.033	0.0441	
CTX-T-CRO-TE-TMP	60	54	23	12	8.94	54.03	< 0.0001	
T-TE-TMP	82	73	55	29	7.31	56.88	< 0.0001	
AX-CTX-T-CRO-TE-CL-AM-L	69	62	34	18	7.23	55.01	< 0.0001	
AX-CTX-T-C-CRO-S-TE-CL-TMP- AM-L	13	12	11	6	6.05	3.44	0.076	
AX-CTX-T-CRO-S-TE-CL-TMP-AM-L	39	35	34	18	2.38	8.99	0.0026	
df = 1 for each resistance pattern								

^{*}Abbreviations: Number (N), df (degree of freedom), Ampicillin (AM), Amoxicillin (AX), Amikacin (AK), Gentamicin (CN), Streptomycin (S), Trimethoprim (TMP), Lincomycin (L), Norfloxacin (NOR), Ciprofloxacin (CIP), Nalidixic Acid (NA), Chloramphenicol (C), Cefotaxime (CTX), Ceftriaxone (CRO), cephalexin (CL), Tetracyclin (TE) and oxytetracyclin (T).

Discussion

Present results show increase dissemination of integrons (37%) among commonsal E. coli isolates this may be because of the widespread of horizontal transfer of mobile elements such as plasmids and transposones that associated with integons between commensal E. coli isolates and *Enterobacteriaceae*. In the same line, Lee et al. (2006) suggest that class 1 integrons harbored by commensal E. coli isolates may be acquired from other pathogenic or commensal bacteria by horizontal transfer of R-plasmids carrying class 1 integrons and that agreement with Sunde (2005), that found rife integrons could be a part of successful plasmids or transposons with a wide, perhaps global dissemination (1,10). In agreement with other studies, this study noted that class 1 integrons were common in E. coli isolates (11,12,13,15). In additional, present study and many other studies demonstrated high resistance of commonsal E. coli to antimicrobials that commonly used for chemotherapy as amoxicillin, ampicillin, cephalexin, cefotaxime and tetracyclines (2,28). Current study similar to study in Mexico that showed resistance to Ampicillin is 100% of commonsal E. coli also like with studies of Marshall et al., (2011) and Li et al., (2014) who showed high resistances of gut flora to streptomycin, nalidixic acid, tetracycline and amoxicillin that may be due to the extensive and long-term use of these antibiotics in humans and livestock (3,9). In additional most recent researches suggest increase and spread antibiotics resistance related to horizontal transfer of resistance genes such as integrons among bacterial populations. Transport resistance genes from commonsal E. coli to pathogenic bacteria represents a potential risk to public health (28,29). Marshall et al., (2009) elucidating that transfer of antibiotic resistance genes from commensals to pathogens depends on the density of donor and recipient cells, the availability of a transfer mechanism, nutrition, and selective pressures and regarded the intestinal environment is considered optimal (28). Low resistance for aminoglycosides (amikacin and gentamicin) among commonsal E. coli isolates recorded in this study agreement with results of Phongpaichit and his coworker (2008) but this results unlike results of Yang et al., (2009) in Taiwan that showed high resistance rates in fecal

strains of E. coli were observed for streptomycin (52%), ampicillin (50.2%), trimethoprim (47.6%) and chloramphenicol (33.8%) (2,18). However the various percentages of resistance in different parts of the world are due to differences in the prevalence of antibiotic consumption in each country (32).

Higher percentage of resistance to some antimicrobial agents (aminoglycoside, cephalosporines, quinolones, and beta-lactam agents) were observed among integron-positive strains with respect to integron-negative strains. The fact could be explained by the presence of resistance genes of these antibiotics in the conserved or variable region of integrons, or by the inclusion of resistance genes in the same mobile elements that carry integrons (17,3). Significant correlated between *IntI1* gene positive isolates and resistance to cefotaxime, tetracycline and trimethoprim (P < 0.05) also determent in results of Li and his coworkers in china that also detected clear association between *IntI1* gene and resistance to nalidixic acid and ampicillin/sulbactam (3). Other study conducted in Taiwan has shown that fecal *E. coli* isolates have high levels antibiotic resistance gene cassette containing class 1 integron (18).

All isolates in present study are multiple drug resistance isolates and are resist to at least four antibiotics but integron positive isolates are resist to at least six antibiotics this may be related to present many resistance genes associated with integrons or may be due to present more than one integron in the same isolate and each one carry out different resistance genes for different antibiotics. Current results agreement with study of Phongpaichit et al., (2008) who found multiple drug resistance was more frequent in integron-positive isolates (89%) than those in integron-negative E. coli (57%) (2). Moreover study of Dureja and his coworker (2014) showed the class 1 integron positive isolates contain eight different resistance gene cassettes in five different combinations, namely dfrA12-orfF-aadA2, dfrA1- aadA1, dfrA17-aadA5, dfrA5 and dfrA7 and this similar to those observed earlier by Karczmarczyk et al (2011) that remember class 1 and class 2 integrons in the collection were found to contain trimethoprim (dfr) and streptomycin (aad) resistanceencoding genes, which are frequently reported in E. coli isolates recovered from various sources, including human, animal, and environmental samples (15,20). The presence of more than one gene most integrons positive samples support the literature cassettes in suggesting that since 1990 there is a prevalence of class 1 integrons carrying multiple resistant gene cassettes (15,21). These data indicate that human fecal E. coli is a reservoir of antibiotic-resistant genes that poses a significant risk of the spread of microbial resistance in the community. In this study *IntI1* gene positive isolates significantly associated with most detected resistance patterns this may be due to high frequency of integron integrated with resistance gene cassettes for multiple class 1 that antibiotics resistance also class 1 integrons, the 5' conserved region encodes a site-specific recombinase (integrase, intII) and a strong promoter or promoters that ensure expression of the integrated resistance gene cassettes for multiple antibiotics in the same *IntI* positive isolates (2). This high frequency of multidrug resistance among *int11*-positive isolates supports the hypothesis of an association between the presence of class 1 integrons and emerging multidrug resistance in E. coli (22).

finally, numerous studies have shown that integrons play a considerable role in the dissemination of MDR in clinical isolates and environmental samples. Fewer studies have aimed at investigating the role of integrons in the propagation and maintenance of MDR phenotypes in non-clinical commensal isolates from humans. There are probably at least two reasons for the lack of studies investigating integrons in commensal isolates from healthy people. One reason may be due to the lack of availability of samples. While bacteria are routinely obtained from hospitalized patients in order to identify and treat infections, healthy people may be far less likely to voluntarily submit to a rectal swab or fecal sample. Perhaps a second reason is that studying bacteria isolated from clinical samples and from individuals with infections can easily seem like a more pressing matter, and seem more vital to understanding the problem of the rapid spread of multidrug-resistance (23).

Conclusion

High prevalence of integron class 1 and antibiotic resistance that produced significant correlation between multiple drug resistance and integron class 1 among *E. coli* flora that may be act as reservoir of antibiotic-resistant genes that has a significant risk for spread of antibiotic resistance to pathogenic or commonsal bacteria in the community.

References

- 1. Lee, J. C., Kang, H. Y., Oh, J. Y., Jeong, J. H., Kim, J., Seol, S. Y., Cho, D. T. and Lee, Y. C., (2006). Antimicrobial Resistance and Integrons Found in commensal *Escherichia coli* Isolates from healthy humans. Journal of Bacteriology and Virology. 36 (3): 133 139.
- **2.** Phongpaichit, S., Wuttananupan, K. and Samasanti, W., (2008). Class 1 integrons and multidrug resistance among *Escherichia coli* isolates from human stools. 39 (2): 279-287.
- 3. Li, B., Zhao, Z., Wang, M., Huang, X., Pan, Y. and Cao, Y., (2014). Antimicrobial resistance and integrons of commensal *Escherichia coli* strains from healthy humans in China. Journal of Chemotherapy. 26(3): 190-192.
- **4. El-Shennawy, G. A., (2011).** Detection of class 1 integrons mediated antibiotic resistance among commensal *Escherichia coli*. Egyptian Journal of Medical Microbiology. 20 (1): 1-10.
- **5.** Oluyege, A. O., Ojo-Bola, O. and Oludada, O. E., (2015). Carriage of antibiotic resistant commensal *Escherichia coli* in infants below 5 months in Ado-Ekiti. Int.J.Curr.Microbiol.App.Sci. 4(5): 1096-1102.
- **6.** Sepp, E., Stsepetova, J., Lõivukene, K., Truusalu, K., Kõljalg, S., Naaber, P. and Mikelsaar, M., (2009). The occurrence of antimicrobial resistance and class 1 integrons among commensal *Escherichia coli* isolates from infants and elderly persons. Annals of Clinical Microbiology and Antimicrobials . 8 (34): 1-6.
- 7. Cocchi, S., Grasselli, E., Gutacker, M., Benagli, C., Convert, M. & Piffaretti J., (2007). Distribution and characterization of integrons in *Escherichia coli* strains of animal and human origin. FEMS Immunol Med Microbiol. 50: 126–132.

- 8. Su, J., Shi, L., Yang, L., Xiao, Z., Li, X. and Yamasaki, S., (2006). Analysis of integrons in clinical isolates of *Escherichia coli* in China during the last six years. FEMS Microbiol Lett. 254(1): 75–80.
- **9.** Marshall, B. M. and Levy, S. B. (2011). Food animals and antimicrobials: impacts on human health. Clin Microbiol Rev. 24:718–33.
- **10.Sunde, M., (2005).** Prevalence and characterization of class 1 and class 2 integrons in *Escherichia coli* isolated from meat and meat products of Norwegian origin. Journal of Antimicrobial Chemotherapy. 56: 1019–1024
- **11.Reyes, A., Bello, H., Dominguez, M., Mella, S., Zemelman, R., Gonzalez, G., (2003).** Prevalence and types of class 1 integrons in aminoglycoside-resistant *Enterobacteriaceae* from several Chilean hospitals. J Antimicrob Chemother. 51: 317-21.
- **12.**Skurnik, D., Menac'h, A. L., Zurakowski, D., Mazel, D., Courvalin, P., Denamur, E., Andremont, A. and Ruimy, R., (2005). Integron-associated antibiotic resistance and phylogenetic grouping of *Escherichia coli* isolates from healthy subjects free of recent antibiotic exposure. Antimicrobial agents and chemotherapy. 49(7): 3062–3065.
- **13.Mathai, E., Grape, M., Kronvall, G.** (2004). Integrons and multidrug resistance among *Escherichia coli* causing community-acquired urinary tract infection in southern India. APMIS.112:159-64.
- **14.Van Essen-Zandbergen, A., Smith, H., Veldman, K., Mevius, D.,** (2007). Occurrence and characteristics of class 1, 2 and 3 integrons in *Escherichia coli*, *Salmonella spp* and *Campylobacter spp*. in the Netherlands. J Antimicrob Chemother. 59: 746–750.
- **15.Dureja, C., Mahajan, S. and Raychaudhuri, S. (2014):** Phylogenetic distribution and prevalence of genes encoding class i integrons and CTX-M-15 extended-spectrum b-lactamases in *Escherichia coli* isolates from healthy humans in Chandigarh, India. Plos One. (9): 1-6.

- **16.Barlow, R. S. and Gobius, K. S. (2006).** Diverse class 2 integrons in bacteria from beef cattle sources. J Antimicrob Chemother. 58:1133–1138
- 17. Vinue, L., Sa'enz, Y., Somalo, S., Escudero, E., Moreno, M. A., Ruiz-Larrea, F., Torres, C., (2008). Prevalence and diversity of integrons and associated resistance genes in faecal *Escherichia coli* isolates of healthy humans in Spain. J Antimicrob Chemother. 62:934—
- **18.Yang, C. M., Lin, M. F., Lin, C. H., Huang, Y. T., Hsu, C. T., Liou, M. L. (2009).** Characterization of antimicrobial resistance patterns and integrons in human fecal *Escherichia coli* in Taiwan. Jpn J Infect Dis. 62(3):177-81.
- **19.Navidinia, M., Peerayeh, S. N., Fallah, F., Bakhshi, B. and Sajadinia, R. S. (2014).** Phylogenetic grouping and pathotypic comparison of urine and fecal *Escherichia coli* isolates from children with urinary tract infection. Braz J Microbiol. 45(2): 509-14.
- **20.**Karczmarczyk, M., Walsh, C., Slowey, R., Nola Leonard, N. and Fanning, S. (2011). Molecular characterization of multidrugresistant *Escherichia coli* isolates from Irish Cattle Farms Appl Environ Microbiol. 77(20): 7121–7127.
- 21.Kang, H.Y., Jeong, Y.S., Oh, J. Y, Tae, S. H., Choi, C. H, Moon D. C., Lee W.K., Lee, Y. C., Seo, S. Y., Cho, D. T. and Lee, J.C. (2005). Characterization of antimicrobial resistance and class 1 integrons found in *Escherichia coli* isolates from humans and animals in Korea. J Antimicrob Chemother. 55: 639–644.
- **22.Yu, T., Zhang, J., Jiang, X.,Wu, J., Dai, Z., Wu, Z., Liang, Y., Wang, X., (2016).** Characterization and horizontal transfer of class 1 integrons in *Escherichia coli* isolates from cooked meat products. Infect Dev Ctries. 10(1):068-073.
- **23.Avila, A. L., (2013):** Prevalence and characterization of integrons in multidrug-resistant non-clinical enteric bacterial isolates. PHD thesis, California State University, Sacramento. 1-96.
- 24.Kazemnia, A., Ahmadi, M. and Dilmaghani, M. (2014). Antibiotic resistance pattern of different *Escherichia coli* phylogenetic groups

- isolated from human urinary tract infection and avian coli bacillosis. Iranian Biomedical Journal. 18 (4): 219-224.
- **25.Clinical and Laboratory Standards Institute (CLSI) (2011).**Performance standards for antimicrobial susceptibility testing, 21th informational supplement Approved standard M100-S21. Wayne, PA: Clinical and Laboratory Standards Institute.
- **26.Kargar, M., Mohammadalipour, Z., Doosti, A., Lorzadeh, S., Japoni-Nejad, A., (2014).** High prevalence of class 1 to 3 integrons among multidrug-resistant diarrheagenic *Escherichia coli* in Southwest of Iran. Osong Public Health Res Perspect. 5(4): 193-198.
- **27.Marshall, B. M., Ochieng, D. J., and Levy, S. B.,** (2009). Commensals: Underappreciated reservoir of antibiotic resistance Microbe. 4:231-38.
- 28.Hsu, S. C., Chiu, T. H., Pang, J. C., Hsuan-Yuan, C. H., Chang, G. N., and Tsen, H. Y. (2006). Characterisation of antimicrobial resistance patterns and class 1 integrons among *Escherichia coli* and *Salmo*nella enterica serovar Choleraesuis strains isolated from humans and swine in Taiwan. nt J Antimicrob Agents. 27(5):383-391.
- **29.Mendonça, N., Figueiredo, R., Mendes, C., Card, R. M., Anjum, M. F. and Silva, G. J. (2016).** Microarray evaluation of antimicrobial resistance and virulence of *Escherichia coli* Isolates from portuguese poultry. Antibiotics. 5 (4):1-9.
- **30.Bailey, J. K., Pinyon, J. L., Anantham, S. and Hall, R.M., (2010):** Commensal *Escherichia coli* of healthy humans: a reservoir for antibiotic-resistance determinants. Journal of Medical Microbiology. 48 (9).
- **31.Kheiri, R. and Akhtari, L. (2016).** Antimicrobial resistance and integron gene cassette arrays in commensal *Escherichia coli* from human and animal sources in IRI. Gut Pathog. 8:40. 1-10
- **32.World Health Organization,**(2014). Antimicrobial Resistance Global Report on Surveillance, Geneva, Switzerland.