was not associated with a positive test (19% vs 20%, p=0.834), but for US-born patients, having a history of travel was associated with a positive test (33% vs 14%, p=0.039). For the Ss positive patients, 34% had a HTLV-I/II test, 48% had at least one stool test, and 76% were given treatment.

Conclusion. There is a significant seroprevalence of Ss in our transplant candidate population, both non-foreign and foreign-born, prompting the indication for universal screening at our facility.

Disclosures. All Authors: No reported disclosures

1387. Impact of Cytomegalovirus Prophylaxis on Healthcare Resource Use and Costs among Kidney Transplant Recipients: A United States Renal Data System-Medicare Linked Database Study

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Session: P-79. Transplant: Studies of Pre-transplant Screening and Evaluation

Background. Cytomegalovirus (CMV) management requires a balance between reducing the risk of CMV infection and avoiding anti-viral toxicities. Limited information is available on the impact of CMV prophylaxis on the healthcare resource use (HCRU) and costs among adult kidney transplant recipients (KTRs) in the United States. Therefore, we examined HCRU and cost associated with CMV prophylaxis stratified by the CMV risk categories among KTRs at 1-year post-KT.

Methods. We identified a cohort of 22,918 adults first-time KTRs during 2011–2017 using the US Renal Data System registry-linked Medicare data. Additional inclusion criteria were to have continuous coverage in Medicare Part A & B for ≥ 6-month pre- and ≥ 12-month post KT and Medicare Part D for ≥12-month post-KT. CMV prophylaxis was confirmed as ≥ 1 prescription fill for valacyclovir/(val)ganciclovir prophylaxis doses within 28 days post-KT.

Results. CMV prophylaxis was utilized in 86%, 82%, and 32% of high, intermediate, and low-risk KTRs with an average cost of prophylaxis per KTRs of \$16,241, \$9481, and \$8,648, respectively. In no prophylaxis groups, valganciclovir was utilized in 52%, 34%, and 36% of KTRs (as either pre-emptive or deferred therapy) with an average cost of \$6,719, \$2,722, and \$431 among high, intermediate, and low-risk KTRs, respectively. Among high-risk KTRs, CMV prophylaxis group had a significantly higher prescription drug cost (\$26,060 vs. \$13,433) but a lower average direct healthcare medical cost (\$84,914 vs. \$101,268), mainly due to lower all-cause hospitalization cost (\$56,758 vs. \$69,852) (Table 1). CMV prophylaxis group had lower rates of all-cause rehospitalization, and CMV-and opportunistic infection (OIs)-related hospitalization compared to no prophylaxis (Table 2). In high-risk KTRs, nearly 32% had myelosuppressive events-related hospitalization, and 15% filled granulocyte colony-stimulating factors with an average cost of \$4,695 per treated KTR.

Conclusion. CMV prophylaxis had a higher cost of medications but had a lower medical cost with including all-cause and CMV-related hospitalizations. Myelosuppressive events were frequent and resource-intensive especially in high and intermediate-risk KTRs.

Outcomes	CMV Prophylaxis Use Stratified by CMV Risk Strata												
	High (D	+/R-) (N=3,520)	Intermedia	te (R+) (N=16,957	0	Low (D-/R-) (N=2,441)							
	Yes	No	Sig. *	Yes	No	Sig.*	Yes	No	Sig.				
Sample Size	3,022 (85.6%)	498 (14.1%)		13,975 (82.4%)	2,982 (17.6%)		784 (32.1%)	1,657 (67.9%)					
Healthcare Costs													
Medical (Inpatient, ER, Out, SNF, HH)	\$84,914 (67,811)	\$101,268 (90,071)	<0.001	\$79,888 (59,331)	\$92,149 (89,836)	< 0.001	\$82,501 (54,605)	\$80,046 (64,800)					
Inpatient hospitalization	\$56,758 (54,035)	\$69,852 (76,964)		\$52,949 (46,882)	\$62,997 (73,548)			\$52,562 (49,298)					
CMV-related	\$3,908(18,843)	\$7,183 (32,330)	0.028	\$933 (8,208)		< 0.001	383.114 (4,883.3)	542.550					
Of-related	\$13,282 (35,399)	\$21,446 (49,708)	0.001	\$12,102 (31,496)	\$18,309 (48,703)	< 0.001	\$10,455 (26,502)	\$11,998 (30,524)					
Myelosupression-related	\$9,932 (23,169)	\$14,043 (34,186)	0.010	\$8,011 (20,689)	\$10,436 (33,019)	0.000	8089.955	\$6,614 (18,333)	0.05				
Emergency Room (ER) Visit	\$656 (1,254)	\$633 (1,400)		602.347 (1,233.7)	\$564 (1,139)	0.097	627.814 (1,904.6)	\$643 (1,526)					
CMV-related	\$7(126)	\$17 (193)		\$2 (75)	\$2 (55)		\$0(0)	\$8 (282)					
Of-related	\$32 (239)	\$22 (141)		\$27 (223)	\$24 (179)		\$21 (219)	\$25 (221)					
Myelosupression-related	\$41(258)	\$52 (324)		\$26 (236)	\$28 (267)		\$26 (206)	\$19 (171)					
Outpatient	\$9,787 (18,019)	\$9,557 (11,894)		\$8,881 (17,965)	\$9,494 (25,110)		\$10,226 (18,854)	\$9,254 (24,902)					
Office-visit	\$9,992 (9,939)	\$11,199 (10,196)	0.012	9800.399	\$10,111 (10,218)		\$9,178 (7,546)	\$9,687 (8,176)					
Durable Medical Equipment (DME)	\$5,099 (4,771)	\$4,917 (4,335)		5392.172	\$5,479 (6,421)		\$6,013 (5,605)	\$5,612 (5,364)	0.08				
Home-Health (HH)	\$1,532 (2,814)	\$2,044 (3,507)	0.002	\$1,602 (3,066)	\$1,800 (3,265)	0.002	\$1,513 (2,892)	\$1,308 (2,605)	0.09				
Skilled Nursing Facility (SNF)	\$1,080 (5,773)	\$3,050 (11,829)	< 0.001	\$651 (4,315)	\$1,655 (7,127)	< 0.001		\$958 (5,056)					
All Pharmacy Claims		\$13,433 (18,215)		\$20,085 (24,410)	\$10,772 (22,060)		\$18,237 (21,350)	\$8,990 (25,552)	<0.00				
CMV Medications	\$16,241 (11,235)		< 0.001	\$9,481 (7,103)	\$2,768 (5,836)	< 0.001	\$8,649 (6,638)	\$475 (2,879)	< 0.00				
Acyelovir	\$2 (32)	\$3 (34)		\$2.7 (51)	\$17 (146)	< 0.001	\$1 (18)	\$23 (329)	0.000				
Valacyclovir	\$1.6(21)	\$3 (34)		\$2 (30)	24 (134)	< 0.001	\$2 (21)	\$17 (77)	< 0.00				
Gancielovir	\$24 (380)	\$68 (835)		\$1 (40)	\$4 (90)		\$0(0)	\$0 (9)					
Valganciclovir	\$16,213 (11,208)		< 0.001	\$9,473 (7,099)	\$2,722 (5,843)	< 0.001	\$8,646 (6,639)	\$432 (2,856)	<0.00				
GCSF related	\$651 (3,136)	\$726 (2,985)		\$356 (3,002)	\$215 (1,558)	0.000	\$203 (1,405)	\$205 (2,430)					
Fotal Cost Medical and Pharmacy	\$110,974 (73,100)	\$114,701 (93,405)		\$99,973 (66,733)	\$102,921 (95,034)		\$100,738 (60,733)	\$89,036 (72,334)	<0.00				

Notes: "All cost data were adjusted to 2019 US dollars using the consumer price index." "Significant differences between the groups were determined by t-tests or analysis of variance (ANOVA) for continous variables

CAV - extraogative (26%: garmlooste colors) standard of the colors and CAVI - servopositive isches you do not and CAVI - servopositive inches you can be colored as a color of the color of

HealthCare Resource Use	CMV Prophylaxis Use Stratified by CMV Risk Strata												
	High (D	+/R-) (N=3,520)		Intermedia	te (R+) (N=16,9	57)	Low (D-/R-) (N=2,441)						
	Yes	No	Sig.*	Yes	No	Sig.*	Yes	No	Sig.*				
	3,022 (85.6%)	498 (14.1%)		13,975 (82.4%)	2,982 (17.6%)		784 (32.1%)	1,657 (67.9%)					
All cause hospitalization													
Inpatient hospitalization	3,022 (100.0%)	498 (100.0%)		13,972	2,980 (99.9%)		784 (100.0%)	1,655 (99.9%)					
CMV-related	432 (14.3%)	105 (21.1%)	< 0.001	530 (3.8%)	196 (6.6%)	< 0.001	10 (1.3%)	19 (1.1%)					
OI-related	997 (33.0%)	220 (44.2%)	< 0.001	4,471 (32.0%)	1,079 (36.2%)	< 0.001	225 (28.7%)	502 (30.3%)					
Myelosupression-related	961 (31.8%)	175 (35.1%)		3,548 (25.4%)	733 (24.6%)		205 (26.1%)	374 (22.6%)	0.052				
Re-hospitalization	1,846 (61.1%)	337 (67.7%)	0.005	7,833 (56.1%)	1,775 (59.5%)	0.001	464 (59.2%)	900 (54.3%)	0.024				
Length of Stay, mean (SD)	19.7 (22.8)	27.8 (33.7)	< 0.001	17.5 (20.0)	22.1 (28.4)	< 0.001	17.5 (18.2)	18.0 (21.3)					
Emergency Room Visit	1,557 (51.5%)	243 (48.8%)		6,975 (49.9%)	1,422 (47.7%)	0.027	386 (49.2%)	846 (51.1%)					
CMV-related	19 (0.6%)	8 (1.6%)	0.020	23 (0.2%)	5 (0.2%)		0 (0.0%)	2 (0.1%)					
OI-related	101 (3.3%)	18 (3.6%)		440 (3.1%)	91 (3.1%)		21 (2.7%)	44 (2.7%)					
Myelosupression-related	114 (3.8%)	22 (4.4%)		329 (2.4%)	71 (2.4%)		18 (2.3%)	32 (1.9%)					
# of office visits, mean (SD)	82.541 (73.8)	92.237 (82.9)	0.014	78.653 (67.2)	81.483 (79.9)	0.071	72.870 (61.7)	81.215 (68.8)	0.003				
Outpatient	3,010 (99.6%)	497 (99.8%)		13,923 (99.6%)	2,970 (99.6%)		781 (99.6%)	1,649 (99.5%)					
Durable Medical Equipment	2,643 (87.5%)	433 (86.9%)		12,395 (88.7%)	2,496 (83.7%)	< 0.001	689 (87.9%)	1,501 (90.6%)	0.040				
Home-Health	1,190 (39.4%)	233 (46.8%)	0.002	5,285 (37.8%)	1,207 (40.5%)	0.007	292 (37.2%)	571 (34.5%)					
Skilled Nursing Facility	173 (5.7%)	71 (14.3%)	< 0.001	580 (4.2%)	247 (8.3%)	< 0.001	33 (4.2%)	103 (6.2%)	0.044				
				13,975									
CMV Medication	3,022 (100.0%)	275 (55.2%)	< 0.001	(100.0%)	1,529 (51.3%)	< 0.001	784 (100.0%)	603 (36.4%)	< 0.00				
Acyclovir	45 (1.5%)	22 (4.4%)	< 0.001	224 (1.6%)	463 (15.5%)	< 0.001	11 (1.4%)	404 (24.4%)	< 0.00				
Valacyclovir	51 (1.7%)	9 (1.8%)		312 (2.2%)	203 (6.8%)	< 0.001	19 (2.4%)	141 (8.5%)	< 0.00				
Ganciclovir	24 (0.8%)	6 (1.2%)		16 (0.1%)	9 (0.3%)	0.016	0 (0.0%)	2 (0.1%)					
Valganciclovir	3,022 (100.0%)	260 (52.2%)	< 0.001	(100.0%)	1,021 (34.2%)	< 0.001	784 (100.0%)	93 (5.6%)	< 0.00				
G-CSF	448 (14,8%)	77 (15.5%)		1.161 (8.3%)	162 (5.4%)	<0.001	58 (7.4%)	58 (3.5%)	< 0.00				

[&]quot;Significant differences between the groups were determined by t-tests or analysis of variance (ANOVA) for continous or chi-square tests for categorical variables.

Disclosures. Amit D. Raval, PhD, Merck and Co., Inc. (Employee) Yuexin Tang, PhD, JnJ (Other Financial or Material Support, Spouse's employment)Merck & Co., Inc. (Employee, Shareholder)

1388. Epidemiology and Treatment Outcomes of Nontuberculous Mycobacterial Infections at a Community Teaching Hospital in the Southeastern United States Y. Vivian Tsai, PharmD¹; Caroline Derrick, PharmD²; Ismaeel Yunusa, PharmD, PhD³; Sharon Weissman, MD⁴; Majdi N. Al-hasan, MD⁴; Julie Ann Justo, PharmD, MS, BCPS-AQ ID⁴; P. Brandon Bookstaver, Pharm D³; ¹Prisma Health Richland - University of South Carolina, Columbia, South Carolina; ²University of South Carolina School of Medicine, Columbia, South Carolina; ³University of South Carolina Columbia, South Carolina; ⁴University of South Carolina, Columbia, SC

Session: P-80. Tuberculosis and other Mycobacterial Infections

Background. Gaps in evidence concerning the epidemiology of nontuberculous mycobacterial (NTM) organisms and their associated treatment outcomes are evident in the literature. The aim of this study was to describe NTM species distribution and susceptibility profile and associated treatment outcomes among adult patients at a tertiary referral hospital in the Southeastern United States.

Methods. A retrospective cohort study of adult patients with NTM infections from January 1, 2010 to June 30, 2020 was performed. Included patients had a positive culture for NTM species and clinical suspicion of infection. Patients were excluded if they had concurrent positive culture for M. tuberculosis (MTB) or monomicrobial culture for M. gordonae. Study endpoints included predictors for favorable treatment outcome, species distribution, and susceptibility at baseline. Favorable treatment outcome was defined as physician-guided cessation of therapy due to clinical improvement. Univariate followed by multivariate regression analysis was used to analyze favorable predictors.

Results. A total of 250 and 78 patients were included in microbiologic and outcomes cohorts, respectively. Among treated patients, 47 (60%) had a favorable treatment outcome. The outcomes cohort consisted primarily of non-Hispanic Caucasians (71%) with pulmonary infection (67%). The most common isolates observed were *Mycobacterium avium complex* (MAC) (67%) and *M. abscessus* (18%). Being self-pay, underweight, history of MTB treatment, and concurrent asthma were more common in those with unfavorable treatment outcomes. The significant favorable predictors included antibiotic change not due to escalation or de-escalation of therapy and private insurance. Among MAC isolates, clarithromycin and amikacin were highly susceptible; however, *M. abscessus* has reduced susceptibility to first-line agents such as amikacin, clarithromycin, and cefoxitin (Table 1).

Table 1. Baseline Susceptibility

	Ciprofloxacin	Moxifloxacin	Clarithromycin	Linezolid	Amikacin	Tobramycin	SXT	Cefoxitin	Imipenem	Doxycycline	Minocycline	Tigecycline	Rifampin	Rifabutin	Ethambutol
# of isolates	11	53	53	53	52		2	1	1	1	1	1		2	3
MAC	0	96.2	98	77.4	96.2		NA	0	0	NA	0	NA		50	0
# of isolates			2											2	
M. kansasii			100											100	
# of isolates	3	3	5	3	5		4	1	1	3	1		3	1	3
Other Slow Growers*	100	100	80	100	100		75	100	100	66.7	100		100	100	66.
# of isolates	44	44	44	44	44		44	44	41	44	43	27			
M. abscessus	0	0	34.1	43.2	86.4		0	13.6	0	0	0	100			
# of isolates	5	5	5	5		5	4	5	5	5	5	4			
M. chelonae	0	0	100	100		100	0	0	0	0	0	100			
# of isolates	14	14	12	14	14		14	14	14	14	14	9	-		
M. fortuitum	100	100	8.3	100	100		85.7	14.3	57.1	28.6	42.9	100			
# of isolates	8	8	8	8	8		8	8	8	8	7	6			
Other Rapid Growers**	62.5	87.5	62.5	87.5	87.5		87.5	25	78	50	57.1	100			

Conclusion. Considering the long incubation time, knowledge of prevalence, antimicrobial susceptibility patterns, and outcomes could guide empirical antimicrobial selection for NTM infections. This is particularly useful for M. abscessus infections where most isolates carry significant resistance to one or more first-line agents.

Disclosures. Julie Ann Justo, PharmD, MS, BCPS-AQ ID, bioMerieux (Speaker's Bureau)Merck & Co. (Advisor or Review Panel member)Therapeutic Research Center (Speaker's Bureau)Vaxart (Shareholder) P. Brandon Bookstaver, Pharm D, ALK Abello, Inc. (Grant/Research Support, Advisor or Review Panel member)Biomerieux (Speaker's Bureau)Kedrion Biopharma (Grant/Research Support, Advisor or Review Panel member)

1389. Nontuberculous Mycobacterial Infections of the Upper Extremity
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Session: P-80. Tuberculosis and other Mycobacterial Infections

Background. Although uncommon, nontuberculous mycobacterial infections (NTMI) of the upper extremity cause significant morbidity based on their natural history, delay in diagnosis, prolonged duration of antimicrobial therapy often combined

ADV cytomegalovirus; D+/R- = CMV-seropositive kidney donor and CMV-seronegative recipient; R+ = CMV-seropositive recipient; D-/R- = CMV-seronegative donor and CMV appropriate (CCM) appropriate