

DRI[™] FENTANYL APPLICATION QuidelOrtho Vitros[®] XT 7600 INTEGRATED SYSTEM, Vitros[®] 5600 INTEGRATED SYSTEM, AND Vitros[®] 4600 CHEMISTRY SYSTEM

Reference No. 10016005, 10016006

This Application is Intended for the Qualitative Determination of Fentanyl in Human Urine.

For Criminal Justice and Forensic Use Only Rx Only

Intended Use	The information provided in this application sheet is intended as a supplement to the package insert. Refer to the package insert for information on intended use, reagent storage, reagent preparation, specimen collection, specimen preparation, specimen storage, quality control, and additional performance data. For package inserts, visit www.quidelortho.com >Resources> MicroTip Partnership Assays (MPA).
Ordering Information	Please place your order with QuidelOrtho. Ordering information available on www.quidelortho.com .
Technical Support Information	Contact QuidelOrtho for technical support. Contact information available on <u>www.quidelortho.com</u> .
	Microgenics Corporation, part of Thermo Fisher Scientific 46500 Kato Road, Fremont, CA 94538 USA

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Unopened reagents are stable until the expiration date when stored at 2 to 8°C. **Reagent Pack** Storage

Reagents stored in UDxx reagent packs onboard the analyzer are stable for 31 days.

Reagents are supplied liquid ready-to-use.

The UDxx reagent pack may be filled up to 15 mL of Reagent A in UDxx/A bottle and 15 mL of Reagent B in UDxx/B bottle, which provides 144 tests.

R1 (mL) in UDxx/A	R2 (mL) in UDxx/B	Tests/pack
15 mL	15 mL	144

If Splitting: The 3 x 18 mL configuration will provide 3.6 UDxx reagent packs with 518 tests, and the 500 mL configuration will provide 33.3 UDxx reagent packs with 4,795 tests.

NOTE: Once the individual UDxx pack number is selected for use during the protocol programming, it is the only UDxx pack number to use for this protocol.

Special Reagent Packs for User Defined Assays

(Please order from QuidelOrtho; not available from Microgenics)

Part Number	Description	Quantity
680 2246	UD01 Packs (Empty)	1 BOX/6PKS
680 2247	UD02 Packs (Empty)	1 BOX/6PKS
680 2248	UD03 Packs (Empty)	1 BOX/6PKS
680 2249	UD04 Packs (Empty)	1 BOX/6PKS
680 2250	UD05 Packs (Empty)	1 BOX/6PKS
680 2251	UD06 Packs (Empty)	1 BOX/6PKS
680 2252	UD07 Packs (Empty)	1 BOX/6PKS
680 2253	UD08 Packs (Empty)	1 BOX/6PKS
680 2254	UD09 Packs (Empty)	1 BOX/6PKS
680 2255	UD10 Packs (Empty)	1 BOX/6PKS
684 4449	UD11 Packs (Empty)	1 BOX/6PKS
684 4448	UD12 Packs (Empty)	1 BOX/6PKS
684 4445	UD13 Packs (Empty)	1 BOX/6PKS
684 4442	UD14 Packs (Empty)	1 BOX/6PKS
684 4447	UD15 Packs (Empty)	1 BOX/6PKS
684 4444	UD16 Packs (Empty)	1 BOX/6PKS
684 4441	UD17 Packs (Empty)	1 BOX/6PKS
684 4446	UD18 Packs (Empty)	1 BOX/6PKS
684 4443	UD19 Packs (Empty)	1 BOX/6PKS
684 4440	UD20 Packs (Empty)	1 BOX/6PKS
680 2256	UDDL1 Packs (Empty)	1 BOX/6PKS
680 2257	UDDL2 Packs (Empty)	1 BOX/6PKS

46500 Kato Road

Results and Data Interpretation	For the Qualitative assay, the Vitros [®] XT 7600 System, Vitros [®] 5600 System and Vitros [®] 4600 System can report both a unit-less Index number and a NEG / POS classification. The assay is calibrated at the cutoff concentration so Index numbers less than the cutoff value are negative and those greater than the cutoff value are positive.		
	NOTE: Qualitative results must only be reported as Negative or Positive. Unit-less Index numbers cannot be used in reportable results.		
Out of Range Error Codes	A very high analyte sample can produce an absorbance outside the Vitros [®] System photometer range and will result in a CB error code. Sample results above the high calibrator should be diluted with negative urine and retested.		
Calibration Frequency	It is recommended that recalibration occur after a reagent pack change, a calibrator lot change, and as required following quality control procedures.		

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DRI Fentanyl Assay-Qualitative QuidelOrtho Vitros[®] XT 7600 System, Vitros[®] 5600 System, and Vitros[®] 4600 System

Full Assay Name: <u>FENTANYL</u>	Version Date: <u>3/14/2019</u>
Short Assay Name: <u>FENT</u>	Fluid Type: <u>Urine</u>
Assay Model Type: <u>2 Point Rate</u>	Template: <u>*2Pt R1-S-R2</u>
Cal Model Type: <u>Linear</u> Calibra	tor Bottles: <u>2</u> Reagent Reps per Cal: <u>2</u>
Reagent Lot Information	Edit Dilution Parameters
-	
On-Board Stability: <u>31</u> Days	Diluent: <u>DATDIL</u> Standard Dilution Factor: <u>1.0</u>
Reagent Lot Num: <u><i>Kit Lot</i></u>	Reflex Dilution: OFF Dilution Factor: 20.0
Shelf Exp. Date: <u>Kit Exp Date</u>	Reduction Factor: <u>1.0</u>
Edit Result Parameters	
Units: <u>%</u>	Number of Qualitative Ranges: 2
Significant Digits: <u>4</u> Precision Digits: <u>0</u>	1. <u>NEGATIVE</u>
User Adjusted Parameters	2. <u>POSITIVE</u> <u>100</u>
Slope: <u>1.0</u> Intercept: <u>0.0</u>	Report Qualitative Result Outside of Range: Yes
CuveTip Exp Time: <u>35</u> Temp Sens : <u>No</u>	Measuring (Reportable) <u>0</u> to <u>10000</u>
	(More Assay Parms) – Edit 2 Pt Rate Additional Parameters
	Initial Abs. Limits: <u>-0.20</u> to <u>3.50</u>
	Second Abs. Limits: <u>-0.20</u> to <u>3.50</u>
	Antigen Excess Factor: <u>9.0</u>

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DRI Fentanyl Assay-Qualitative QuidelOrtho Vitros[®] XT 7600 System, Vitros[®] 5600 System, and Vitros[®] 4600 System, *continued*

		Edit Protocol Parameters			
	Step	Volume	Pack ID	Seconds	Wavelength
1.	Reagent	80 μL	UDxx/A		
2.	Incubation			0.0	
3.	Sample	17.2 μL			
4.	Incubation			304.0	
5.	Reagent	80 μL	UDxx /B		
6.	Incubation			38.0	
7.	Read				340 nm
8.	Incubation			76.0	
9.	Read				340 nm

Edit Calibration Parameters

Bottle #	Dil Factor	Cal Rep Resp Range	Calibrator Lot: <u>Cal Kit lot</u>
1	_1.0		Cal value: <u>0</u>
2	1.0	0.20	Cal Value: <u>100</u>

(More Cal Parms) – Edit Linear or Logit/Log Additional Parameters

Monotonicity: <u>Increase</u>		
Max Resp High: <u>3.00</u>	Min. Resp. High: <u>3.00</u>	Cal Fit Goodness Limit: <u>0.950</u>
Max Resp. Low: <u>-3.00</u>	Min Resp. Low: <u>-3.00</u>	

Edit Triple Read Parameters

	Reportable Conc.	Triple Read Limit
Reportable Min.:	<u>0</u>	[Let it fill in]
Critical Conc.:	2	%
Reportable Max.	<u>10000</u>	<u>8.0</u> %

Comments: N/A

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