SPECIAL 510(k): Device Modification Decision Summary

To: Quidel Corporation **RE:** K131619

This 510(k) submission contains information/data on modifications made to the SUBMITTER'S own Class II device requiring 510(k). The following items are present and acceptable

1. The name and 510(k) number of the SUBMITTER'S previously cleared device:

Trade Name: QuickVue® Influenza A+B test

510(k) number: K092698, K053146, K031899

- Submitter's statement that the INDICATION/INTENDED USE of the modified device as described in its labeling HAS NOT CHANGED along with the proposed labeling which includes instructions for use, package labeling.
- 3. A description of the device **MODIFICATION(S)**. The modification presented in this 510(k) consisted of **expanded reactivity table to include reactivity information for** the H7N9 influenza A virus. The submitter tested the ability of the QuickVue® Influenza A+B test to detect H7N9 influenza A virus. The virus used (A/Anhui/1/2013) was obtained from the Centers for Disease Control and Prevention as non-infectious beta-propiolactone inactivated virus. An LoD study was performed with the A/Anhui/1/2013 influenza strain at concentrations:
 - 7.90x10⁸ EID₅₀/mL
 - 7.90x10⁷ EID₅₀/mL
 - 7.90x10⁶ EID₅₀/mL
 - 3.95x10⁶ EID₅₀/mL

The LoD was determined to be 7.90x10⁶ EID₅₀/mL.

The QuickVue[®] Influenza A+B test package insert has been updated to include the additional analytical reactivity information

- 4. The FUNDAMENTAL SCIENTIFIC TECHNOLOGY of the modified device has not changed.
- 5. **Comparison Information** (similarities and differences) to applicant's legally marketed predicate device including, labeling, intended use, and physical characteristics:

Similarities

Features	QuickVue Influenza A+B test	QuickVue Influenza A+B test
Intended Use	The QuickVue Influenza A+B test	The QuickVue Influenza A+B test
	allows for the rapid, qualitative	allows for the rapid, qualitative
	detection of influenza type A and	detection of influenza type A and
	type B antigens directly from nasal	type B antigens directly from nasal
	swab, nasopharyngeal swab,	swab, nasopharyngeal swab,
	nasal aspirate, and nasal wash	nasal aspirate, and nasal wash
	specimens. The test is intended	specimens. The test is intended
	for use as an aid in the rapid	for use as an aid in the rapid
	differential diagnosis of acute	differential diagnosis of acute
	influenza type A and type B viral	influenza type A and type B viral
	infections. The test is not intended	infections. The test is not intended
	to detect influenza C antigens.	to detect influenza C antigens.

	Negative results should be confirmed by cell culture; they do not preclude influenza virus infection and should not be used as the sole basis for treatment or	Negative results should be confirmed by cell culture; they do not preclude influenza virus infection and should not be used as the sole basis for treatment or
	other management decisions. The test is intended for professional	other management decisions. The test is intended for professional
	and laboratory use.	and laboratory use.
Read Results	Visual	Visual
Specimen Types	Nasal swab, nasopharyngeal swab, nasal aspirate, and nasal wash	Nasal swab, nasopharyngeal swab, nasal aspirate, and nasal wash
Read Result Time	10 minutes	10 minutes
External Controls	Test kit contains Positive and Negative Control swabs	Test kit contains Positive and Negative Control swabs

Differences

The package insert has been updated to include detection of the A/Anhui/1/2013 H7N9 virus in the analytical reactivity information section:

A/Anhui/1/2013 - A - H7N9 - 7.90x10⁶ EID₅₀/mL

Although this test has been shown to detect these 2009 H1N1 and H7N9 viruses cultured from positive human respiratory specimens, the performance characteristics of this device with clinical specimens that are positive for these 2009 H1N1 or H7N9 influenza viruses have not been established. The QuickVue® Influenza A+B Test can distinguish between influenza A and B viruses, but it cannot differentiate influenza A subtypes.

6. Design Control Activities Summary:

- a) Analytical Reactivity Testing was conducted as described in section 3, Device Modifications.
- b) Declaration of Conformity
- A "Declaration of Conformity" statement was submitted for the manufacturing facility and validation activities and signed by the Director of Quality Assurance and the Senior Director of Technical Operations respectively. The statements indicate that;
- 1. The manufacturing facility is in conformance with design control procedure requirements as specified in 21 CFR 820.30 and the records are available for review.
- The validation activities, as required by the risk analysis, for the modification were performed by the designated individuals and the results demonstrated that the predetermined acceptance criteria were met.

In conclusion, based on the results of the analytical reactivity testing the modified labeling is truthful and accurate. The changes do not affect the performance of the test and it is therefore substantially equivalent to the current cleared test.

7. A Truthful and Accurate Statement, a 510(k) Summary, and the Indications for Use Enclosure.

The labeling for this modified subject device has been reviewed to verify that the indication/intended use for the device is unaffected by the modification. In addition, the submitter's description of the particular modification and the comparative information between the modified and unmodified devices demonstrate that the fundamental scientific technology has not changed. On this basis, I recommend the device be determined substantially equivalent to the previously cleared device.