Introduction
Calcium is a ubiquitous eukaryotic signaling molecule that regulates a wide array of cellular processes including fate determination, excitability, motility and adhesion, and regeneration. While there are numerous methods for monitoring calcium activity through both chemical and genetically encoded indicators, there is currently a lack of tools for higher-level measurement such as *in vivo* calcium spike counting. Because developing advanced circuits is very challenging in eukaryotic systems, the William & Mary iGEM team aimed to create a toolkit for inducing, monitoring, and measuring calcium activity in bacteria. This set of foundational tools will allow future synthetic biologists to develop more sophisticated genetic systems for studying calcium.

Objectives
We planned to create three different classes of parts that: induce calcium influx in bacteria; indicate the presence of calcium; and record the number of calcium spikes.

1. **Induce calcium influx in bacteria:**
Channelrhodopsins are light-activated cation channels that, when excited, allow positively charged ions to flow from the extracellular space into the cell. There are two naturally existing channelrhodopsins: channelrhodopsin-1 (ChR1) and channelrhodopsin-2 (ChR2). Additionally, there exist versions, including CatCh, that have been optimized for specific situations.

2. **Indicate the presence of calcium:**
Genetically Encoded Calcium Indicators (GECIs) are fusions of a fluorescent protein and calmodulin. The fluorescent protein is inactive until calmodulin binds to calcium. This induces a conformational change that activates the fluorescent molecule, allowing for the monitoring of changes in calcium levels.

3. **Record the number of calcium spikes:**
A calcium-dependent promoter would activate a counting circuit that, after a specified number of spikes, produces a reporter molecule.

Parts Submitted
- **ChR1** (BBa_K1409000): Channelrhodopsin-1 is a light-gated ion channel that is activated by light of approximately 560nm (yellow light). ChR1 is a non-specific cation channel, allowing the influx of ions such as calcium into cells. 
- **CatCh** (BBa_K1409001): Calcium Relocating Channelrhodopsin (CatCh), is a ChR2 variant with increased permeability to Ca\(^{2+}\).

Conclusions
We successfully created and submitted two new parts: ChR1 (BBa_K1409000) and CatCh (BBa_K1409001). These two channelrhodopsins have been designed to meet the BioBrick RFC10 standard, ensuring that these parts can easily be used by future teams.

Future directions
The next step towards completing a set of tools to enable the study of calcium dynamics in bacteria is to build BioBricks for reporting and measuring calcium activity. We are confident that the introduction of these essential parts will enable future iGEM teams to create genetic circuits for monitoring calcium in bacteria, and eventually eukaryotes.

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**Visualization of transient calcium activity in a developing Xenopus laevis embryo.** Cells are outlined with MemRFP (red) and calcium activity is indicated by GCaMP (green). This is an example of current calcium visualization techniques, but is not capable of recording calcium history. Photo courtesy of Wendy Herbst.

**Restriction digestion confirmation of ChR1 BioBrick clones!**

**The three phases of calcium manipulation in bacteria:**

**Phase I:** Ca\(^{2+}\) concentration in the cell is increased through activation of a calcium-specific ion channel. 

**Phase II:** Increase in calcium concentration activates a genetically encoded calcium indicator (GECI). 

**Phase III:** After a user-defined number of spikes in calcium concentration, transcription of a different fluorescent indicator is activated.